A TREATISE . ~

ON

QUALITATIVE ANALYSIS

ADAPTED FOR USE IN THE LABORATORIES OF COLLEGES AND TECHNICAL INSTITUTES

Ву

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WITH 84 ILLUSTRATIONS

Fourteenth Thousand



J. & A. CHURCHILL
7 GREAT MARLBOROUGH STD

PREFACE

THE aim of this laboratory textbook is to give all the necessary working directions so simply and fully as to reduce to a minimum the necessity of tutorial assistance.

The dimensions of the book have at the same time been kept within reasonable limits by omitting all matter which does not directly bear upon analytical work, and by avoiding lengthy theoretical explanations.

The analytical reactions and methods which are described have been carefully selected; and as they have been frequently worked through from the text by the members of our classes, their accuracy and intelligibility are, we believe, assured.

As a rule the analytical methods have been selected on account of their simplicity and the rapidity and ease of their execution. But occasionally preference has been given to methods which are delicate and trustworthy rather than rapid and simple, and the simpler methods are only recommended as substitutes for these when minute quantities of substance have not to be detected.

It has also been thought advisable to select reactions which are useful in analysis, in preference to those which claim only to be interesting or instructive; and we have not hesitated to adopt methods which are unsuitable for quantitative analysis, if they are rapid and accurate for qualitative purposes.

A glance at the contents on pages vii—xv will show that the book is divided into four main parts and that these are subdivided into eight sections.

· Part I deals with the apparatus and operations which are employed in qualitative analysis.

Part II describes the analytical reactions which are used for the detection of substances, and the methods of applying these reactions.

Part III sets out the procedure for systematically employing these reactions for the detection of the composition of substances, and includes the usual analytical tables.

PREFACE

Part IV contains suggestions for fitting and furnishing the laboratory, together with full lists of the apparatus, reagents and chemicals which are required for analytical works.

The construction and use of the general apparatus required in the laboratory are also described; and suitable methods for preparing solutions of the requisite strength are set out, with the object of securing economy in time and expense. Tables of atomic weights, and of metric and English weights and measures follow at the end of this part.

The arrangement of all the tabular matter across instead of along the page will be found convenient, since it renders unnecessary the turning of the book when the table is being consulted.

Symbolic notation has been freely employed, but any perplexity to beginners which might arise from this source has been avoided by attaching the name of the substance to its symbol or formula, when this is used for the first time in the text. Both names and formulae are also associated in the tables in the eighth section, and it is recommended to have both the name and the chemical formula printed on every bottle-label in the laboratory.

Our acknowledgments for valuable suggestions are due to experienced teachers, and we have found Atack's Chemist's Year-Book most useful in furnishing the many recent chemical and physical constants which appear in the text.

The present edition has been rewritten, recast and enlarged in order to adapt it to modern methods of teaching. The reactions of the metals have been rearranged and placed in the natural order of the metal groups; and among the additions which have been made are the statement of the properties of metals, and many resolutes, mainly for organic substances and the rarer elements:

The introduction of the modern system of standard strengths of the laboratory solutions will be found convenient.

The book is presented in an altered style and size of page which metals it a companion volume to the later editions of our Quantita-

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APPARATUS FOR EACH BENCH-LOCKER

- I Bunsen-burner with tubing (2).
 - 1 Test-tube stand, with twelve holes.
- 12 Test-tubes.
 - 1 Test-tube brush (19).
- 2 Boiling-tubes.
- 2 Round glass plates.
- 2 Porcelain dishes.
- 2 Watch-glasses.
- 1 Four-ounce (120 c.c.) co
- 1 Pestle and mortar.
- 1 Iron tripod-stand.
- 1 Piece of coarse iron wire-gauze.
- 3 Glass funnels, and some cut filter-papers.
- 3 Small beakers.
- 3 Glass rods (15).
- I Piece of platinum-foil.
- 2 Pieces of platinum-wire (14).
- 1 Mouth-blowpipe (II).
- 1 Pipe-clay triangle (fig. 81, 1308).
- 1 Wash-bottle (17).
- . 1 Retort-stand.
 - 1 Wooden filter-stand (fig. 50, 43).
 - 1 Pair of brass crucible tongs.
 - I Small horn or vulcanite spatula.
 - 1 Wicker draining-basket (21).

A fuller description of this apparatus will be found in paragraph 1368. Most of the apparatus may be seen on the bench in Fig. 82 (1360). The reference numbers in brackets refer to paragraphs in which the apparatus is described.

PART I APPARATUS' AND OPERATIONS'

SECTION

DIRECTIONS FOR THE PREPARATION AND USE OF APPARATUS

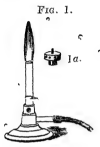
1. General Remarks. A student who is commencing work in a general laboratory will find on the preceding page and in par. 1368 a list of the apparatus which should be contained in his private locker. This apparatus should be looked through, fitted, and cleansed, as is directed in this Section.

In Section VIII. (1369 et seq.) will be found particulars of other apparatus, which may be used by many students in common; also lists of the various reagents, and the methods of preparing them.

For convenience in reference, paragraph numbers are frinted in the text in black type, and in square brackets at the top of each

FLAMES FOR HEATING.

2. The Bunsen-burner (Fig. 1) is generally employed in the laboratory for heating purposes. It is so constructed that coalgas, entering near its base, is mixed with a proper proportion of air before it is burnt.



BUNSEN-BURNER.

The air is drawn in through holes at the lower part of the burner, and the mixture of gas and air is burnt at the top of the upright tube. The oxygen of the air, which is thus mixed with the gas, burns the carbon in the body of the flame. Accordingly the luminosity, which depends upon the existence in the flame of unburnt carbon or carbon compounds, is destroyed.

The Bunsen-flame is therefore mainly useful, because it deposits no particles of soot when it is in contact with any surface. Its high

temperature and non-luminosity also render it very valuable for producing flame-colorations.

The burner should be provided with some means for partly or entirely closing the air-holes when requisite. This is usually affected by turning round a loose perforated ring outside the holes. If the flame is reduced to small dimensions, the supply of air should be partly shut off, else the flame is apt to recede.

When the burner is to be used, it is connected by means of a piece of tightly-fitting rubber tubing, about 5-16ths of an inch (7.5 mm.) in internal diameter, with the tube which supplies gas to the working-bench. The tap is then turned on, and in a few seconds the gas is lighted. The flame should be almost perfectly colourless and non-luminous.

Occasionally the gas will "burn below," that is, at the bottom instead of at the top of the burner. This usually happens when the flame is small and the quantity of air admitted through the holes is too great. It also occurs when the burner is lighted too soon after turning on the gas-supply. A luminous flame may then be seen through the holes near the base of the burner, burning from the small gas-jet inside. The flame at the top of the burner will also be long and somewhat luminous, and will emit a very unpleasant smell, which is due to the incomplete combustion of the gas.

In such a case the flame should be at once extinguished by pinching the rubber tubing close to the burner, and the gas should be relighted after it has escaped for a few seconds

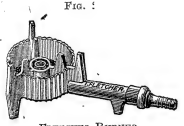
The above statements should be proved by trial, and the effect of partially or entirely closing the air-holes of the burner should be observed.

The Rose-top (Fig. 1a) is employed for diffusing the heat of the flame over a large surface. It is a small perforated metal cap which is placed upon the top of the burner, and yields a circle of flames. The rose-top is put on the burner and removed from it by means of crucible-tongs. It is of course very hot after being used, and should never be handled, or placed upon wood or glass, until it is cool.

3. High Temperature Burners of several forms are in use for special purposes. The Fletcher and Méker burners are arranged

to mix the gas and air in such proportions as to produce "solid flame" when the mixture is burnt. The temperature in all parts of such a flame is the same and is relatively high. These flames may therefore be substituted for the blowpipe flame (FI) for many purposes.

A convenient form of Argand Fletcher-burner for supporting



FLETCHER BURNER.

and heating large vessels is shown in Fig. 2; its flame is compact, and in regulating the size of the flame the proportion of gas to air is automatically adjusted.



TECLU BURNER.

The Teclu Burner (Fig. 3) is a suitable high temperature burner for use on the workingbench. In this burner the air enters through the conical part and its supply is regulated! by rotating the disc (a), and the gas-feed is regulated by the rotation of the screw (b). When the highest temperature is required, the full supply of gas is turned on, and the disc is rotated until a coaring flame is produced.

4. Other Special Burners (5, 6, 7) are . occasionally used instead of gas-burners; but

for general purposes they should only replace the gas-burners when coal-gas cannot be obtained.

5. The Spirit-lamp (Fig. 4) consists of a glass vessel containing methylated spirit, into which dips a cotton wick supported by means of a brass or stoneware wick-holder.

When the lamp is not in use the upper end of the wick should always 😓 covered with the glass cap, in order to prevent evaporation of the spirit. If the spirit is tolerably free from resinous matter, its flame will be non-luminous and will deposit no soot.

6. Vapour Burners, arranged to burn a mixture of air with vaporised methylated spirit or petroleum, are now obtainable, and may be used in



SPIRIT-LAMP.

laboratories where a supply of coal-gas is not available.

A burner similar in form to the Bunsenburner (2) for burning a mixture of vaporised methylated spirit with air, and a metal lamp which furnishes a smokeless non-luminous flame with vaporised petroleum oil, are described in peragraph 1366.

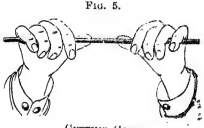
7. An Acetylene Bunsen-burner may also be used, the acetylene gas being inade as required in an automatic apparatus by the action of water on calcium carbide, or a supply of the gas being stored in a small gas-holder.

Such a burner can be used in laboratories which are situated at a distance from gas-works, and a laboratory attendant, can easily perform the duties of gas-maker if the gas is stored.

CUTTING AND BENDING GLASS TUBE AND ROD.

8. Glass Tube or Rod is Cut by laying it upon a flat surface, and making a deep scratch with the edge of a three-cornered file at the point to be cut. The glass is then held with both hands, one on each side of the scratch and close to it, and a gentle pressure is exerted upon the glass as if trying to break it across (Fig. 5). If the file-mark has been made sufficiently deep, the glass will readily break at this point.

The sharp edges of a freshlycut rod or tube should always be at once rounded. This is effected either by holding the end of the tube in the Bunsen-flame or blowpipe-flame until the edges are partly melted (Fig. 8, page 6), or by rubbing off the sharp edge with the face of a file.

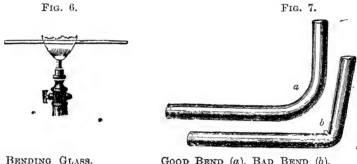


CUTTING GLASS.

Select a length of glass tubing, which is about one-eighth of an inch (3 mm.) in internal diameter. Cut from it one piece about ten inches (25 cm.) and two pieces about five inches (12.5 cm.) long, and carefully round off their sharp freshly-cut edges. Also cut off three pieces of glass rod, respectively seven, six, and three inches (18, 15, and 7.5 cm.) in length. Keep these for future use.

o. Ordinary Glass Tube is Bent by holding it in the upper edge of a common fish-tail gas-flame, so as to heat at least two inches of the glass (Fig. 6). The tube is supported by both hands. one on either side of the flame, and is constantly turned slowly round on its axis, in order to heat all sides equally. As soon as the glass is felt to be soft and pliable, it is taken out of the flame and is at once quickly bent to the required angle. The heated part must not be allowed to touch anything until it is cold. The soot is then removed from it by a cloth or by a piece of paper.

If a bend is properly made, it should be a curve and should not alter the bore of the tube (Fig. 7 a). If a sharp angle is made,



GOOD BEND (a), BAD BEND (b).

the bore will be narrowed (Fig. 7 b), and the bend will not only be unsightly, but weak and liable to break under a small strain.

Bend the longest piece of glass tubing (8) at right angles, so that the shorter part is about two inches (5 cm.) in length. Keep this for future use.

10. Glass rod, Hard glass tube, and Tube with Thick Walls or Small Bore may be bent in the Bunsen-flame, or even in hotter flames including the blowpipe-flame. But these flames are not suitable in shape for bending ordinary glass tube, as they soften the walls too much and cause it to get out of shape.

THE BLOWPIPE, AND ITS USE.

11. The Mouth blowpipe is used for producing a small but very hot flame, which can be made to assume any desired direction. This is effected by blowing a fine stream of air from the mouth through a gas-flame.

The blowpipe is held in the right hand, with its finely pierced tip resting on the edge of the burner, and just inside the flame (Fig. 8). The mouthpiece is then taken between the lips; and after the cheeks have been blown out to their full extent, the air contained in them is forced out through the jet. This produces a small pointed conical flame in the direction of the blast.

The chief difficulty in using the blowpipe properly is experienced in maintaining the blast of air uninterrupted by breathing. Patient trial will, however, remove this difficulty, if it is borne in mind that the cheeks are to be kept constantly inflated with air, and that the air must be forced through the blowpipe by the muscles of the cheeks, and not by the action of the lungs, breathing being meanwhile carried on through the nose, and the mouth being occasionally rapidly replenished from the lungs.









DRAWING OFF GLASS.

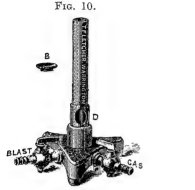
It is frequently necessary to have both hands free while the blowpipe is being used. This may be secured by resting the blowpipe-jet upon the top of the gas-burner and supporting the mouth-piece by the lips alone, as is shown in Fig. 9.

A small flame, made luminous by nearly closing the air-holes of the Bunsen-burner, is better suited for most blowpipe-work than the ordinary non-luminous flame of the burner.

It is important that the hole in the nozzle of the blowpipe should be small and round, else a good flame will not be obtained; and unless the hole is very small it will be found difficult to maintain a continuous blast.

The student should, as soon as possible, acquire dexterity in the use of the mouth-blowpipe, since its flame serves for many of the shorter heating processes of the chemical laboratory. 12. A Blast blowpipe (Fig. 10) which is fed with air from the foot-bellows (Fig. 11), or from a mechanical or water blower, is often advantageously employed for maintaining a high temperature in fusion processes, or for extensive glass-working or glass-blowing.

The special form of Bunsen-blowpipe shown in Fig. 10 is suitable for many purposes. It consists of a somewhat large Bunsen-burner (2), which is furnished with two taps. One of



BUNSEN-BLOWPIPE.



Fig. 11.

FOOT-BELLOWS.

these taps controls the gas-supply, and the other controls the air blast from the bellows.

When a Bunsen-flame is required, the gas is supplied to the burner, and the air-supply is adjusted at the base of the burner by the ring D.

When the blowpipe-flame is to be used, the metal ring B is placed upon the top of the burner so as to reduce the aperture, and the supply of gas and the air-blast are regulated until a flame of the necessary character is obtained.

The Bunsen Gas-blowpipe (1365) will be found better adapted than the above for more elaborate glass-blowing and working.

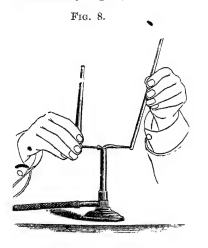
The Herapath Gas-blowpipe (1365) is convenient for smaller heating purposes. It can, if desired, be blown by the mouth. It may be purchased without the foot and the tap, the upright tube being slipped down the tube of a Bunsen-burner.

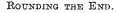
A Petrol, or Spirit, Blowpipe (1366) may be used where coal-gas is not available. It possesses the advantage of being portable and of requiring no bellows or air-blast.

13. Small Ignition-tubes.—One of the pieces of glass tubing, about five inches (13 cm.) in length (8), is drawn out at its middle point by heating it strongly in the blowpipe-flame (Fig. 9, page 6).

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DRAWING OFF GLASS,

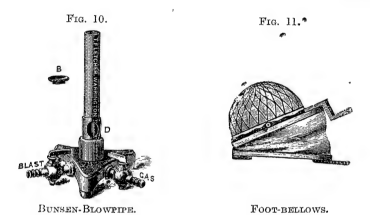
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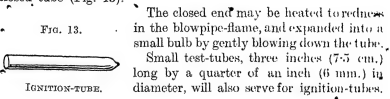
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While the tube is being heated, it is held in both hands, and is constantly turned round upon its long axis in order to secure uniform heating. As soon as the glass is softened, it is gradually drawn out by pulling the ends of the tube in opposite directions:

The narrowed portion of the tube (Fig. 12) is then cut across



(8) at its middle point, the conical part (a) is heated in the blow-pipe-flame, and the narrow portion is drawn off, producing a small closed tube (Fig. 13).



14. Mounted Platinum Wires.—Select a length of platinum wire about as stout as a small pin. Ascertain that the wire does not permanently impart colour to a Bunsen-flame when it is heated to bright redness in the flame. Cut off two pieces of this wire, each about two inches (5 cm.) in length, and proceed to fix them in glass handles in the following way. This will facilitate their being held in the fingers, and will help to prevent them from being lost.

Draw out at its middle point (Fig. 12,) the piece of glass tube (8) about five inches (13 cm.) in length, and cut it across at the middle of the narrow portion. Each of the pieces of tube thus obtained serves for the handle of a wire.

Break off the narrow part of the tube until it extends only about a quarter of an inch (6 mm.) from the shoulder, a (Fig. 12). Insert the end of the platinum wire into this narrow opening. Then hold

Fig. 14.



MOUNTED PLATINUM WIRE.

the end of the tube, containing the wire, in the blowpipe flame, until the glass melts and thickens around the end of the wire, fixing it firmly when cold (Fig. 14).

The free end of the wire may then be rolled round a stout wire into a loop about an eighth of an inch (3 mm.) across.

15. Glass Stirring-rods.—Remove by a file affly small projections from the ends of each of the glass-rods, which were made (8) by cutting some solid glass-rod into lengths of about three, six, and seven inches (7.5, 15 and 18 cm.). Then heat both ends of each rod to redness in the tip of the blowpipe-flame, the rod being meanwhile constantly turned round on its long axis (Fig. 8, page 6). The sharp edges of the glass are thus rounded by fusion. The end of the rod must not be allowed to touch anything until it is cool.

Sometimes a very slender rod is required. This may be obtained by heating a suitable point in an ordinary glass rod with constant rotation in the blowpipe-flame until the glass is well softened, and then drawing it out to the requisite fineness. See Fig. 9 (page 6) and Fig. 12.

Boring Corks.

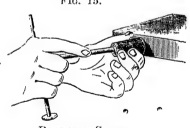
16. A Cork may be Bored, for the insertion of a glass tube, by gradually pushing through it a sharply pointed and slender round file with a constant twisting movement, and then enlarging the hole with the file to the necessary size.

The Cork-borer, which is a brass tube sharpened at one end, is, however, more commonly used.

A borer must be selected which is rather less in size than the

glass tube which is to be inserted into the cork. cork is then pressed against a wooden surface or grasped firmly in the hand, and the perforation is made by gently pushing the borer through it with a constant movement of rotation backwards and forwards upon its axis (Fig. 15).





BORING A CORK.

Caution and practice will enable the student to make a clean straight hole without damaging the surrounding parts of the cork.

In making a single central hole through a cork, it is well to bore from the centre of one end of the cork half-way towards the other end, and then reverse the cork and bore a hole to meet this from the centre of the opposite end. In this way a hole which is central in position throughout the cork is most easily obtained.

A slender round file is used for smoothing the sides of a hole which has been made by a cork-borer, or for slightly enlarging it. Great care must be taken to leave the hole round in shape, and not to enlarge it so much that the glass tube fits loosely.

The cutting edge of the cork-borer is sharpened by rubbing its inner surface with a round file, and then rubbing the outer surface obliquely with the face of a flat or triangular file.

Rubber Corks are perforated in a similar manner by a sharp borer which is either partly filled with water, or is moistened with alcohol or glycerine.

FITTING THE WASH-BOTTLE.

17. The Wash-bottle.—A thin, flat-bottomed, conical flask, about eighteen ounces (500 c.c.) in capacity, and not less than, an inch (2.5 cm.) in diameter in the neck, should be fitted as is

shown in Fig. 16.

Fig. 16.



THE WASH-BOTTLE.

For this purpose a sound cork is selected which is slightly too large to enter the neck of the flask. The cork is softened by rolling it backwards and forwards under the foot with gentle pressure; it must now fit tightly into the neck of the flask.

Two pieces of glass tubing, rather longer than the tubes (a, b), are then bent (9) into the form shown in Fig. 16. Their ends are cut off to the right length, and the sharp edges are rounded by holding them in the Bunsen-flame, or by rubbing them with the face of the file.

Two parallel holes are then bored through the cork, with a proper sized cork-borer (16).

or by means of a round file. The holes must be somewhat smaller than the glass tubes, and must not run into one another or to the outside of the cork. They are smoothed, and slightly enlarged if necessary, by means of the round file.

Into these holes the tubes (a, b) are then gently pushed with a twisting motion. They must enter somewhat stiffly, but without requiring much pressure. If the holes have been carelessly made too large, the tubes may often be made to fit by slipping upon them little pieces of wetted narrow rubber tubing, or by putting the

wetted pieces of rubber tubing into the holes in the cork before the glass tubes are pushed in.

• A Rubber Cork is much more durable than an ordinary cork for this and for most other chemical purposes. It may be purchased with two holes already made, or may be perforated by a sharp well-wetted cork-borer (16), or by a wetted round file. Both the glass tubes and the inside of the holes should be wetted before the tubes are pushed in, since water serves as a lubricant for glass against rubber.

Before the further fitting of the wash-bottle is proceeded with, the cork with the tubes is inserted into the neck of the flask. One tube is then closed with the finger, and air is blown through the other tube into the flask, so as to produce pressure in its interior. The cork is meanwhile carefully watched in order to ascertain whether there is any escape of air. A leakage is as a rule easily heard: but if the outside of the cork is wetted with water, any escape of air-bubbles will be readily seen.

If the cork is air-tight, a piece of rubber tubing, about an inch (2.5 cm.) in length, is pushed upon the upper end of the tube (a). A short jet (c), which has been made by drawing out a piece of glass tubing in the flame (Figs. 9, 12, pages 6, 8), is fitted into the other end of the rubber tube. The narrow opening of the glass jet may be further contracted, if necessary, by holding it in the flame for a short time and slightly fusing the edges of the glass.

If the wash-bottle is to be heated, the neck of the flask should now be bound round with twine like the handle of a cricket bat, or tightly covered with a folded strip of flannel. The neck, when thus protected, can be grasped with comfort even after water has been boiled in the flask and the neck has become heated by steam.

The wash-bottle is now filled about two-thirds with distilled water, and is ready for use. Tap-water should never be kept in the wash-bottle.

A fine stream of water may be obtained from the jet (c), by blowing down the tube (b).

This stream serves for washing precipitates and for other purposes. A larger stream may be obtained by inverting the flask, when the water will flow out from the end of the tube (b), air entering meanwhile by the tube (a): this stream can be made to issue with greater force and speed by blowing down the tube (a).

When hot water is required, the wash-bottle is supported on a tripod-stand upon a piece of coarse iron-wire gauze, and is heated by the Bunsen-flame: or it may be heated by the small Fletcher-burner (Fig. 2, page 3).

^{18.} When Boiling Water, or a Liquid which gives off Vapour or Gas,

is used in the wash-bottle, a special stopper, shown in Fig. 17, should be fitted into the neck of the bottle, in order to prevent the vapour or gas from entering the mouth.

The stopper is perforated with three holes. Through the centre hole is pushed a short glass tube (a), which terminates just above and just below

the stopper.



FITTING FOR HOT-WATER BOTTLE.

The other holes carry the jet-tube, the upper part of which is shown (b), and the blow-tube (c), as in the ordinary washbottle. The lower end of the blow-tube, however, is fitted with a valve, which is formed by slipping over it a piece of rubber tube with a slit cut in it (c) and which is closed below by a piece of glass rod.

While the wash-bottle is being used the tube (a) is closed with the finger, and air is blown in through the valve (c). The valve prevents the air from returning through the blow-tube and introducing vapour or gas into the mouth. When the stream is to be stopped, the finger is removed from the tube (a) and the pressure of air in the interior of the flask is released.

CLEANING APPARATUS.

It is indispensable that all glass and porcelain apparatus should be kept scrupulously clean, and the student should cleanse his set of apparatus as is directed below. In ordinary work the apparatus should be cleansed before it is put away.

19. Test-tube Brush.—This brush is constantly in use for cleansing glass and porcelain apparatus.

The piece of sponge, which is sometimes fastened on the end

of the brush (Fig. 18 a), does not adapt itself well to the bottom of test-tubes and boiling-tubes. A much more efficient end is given to the brush by removing the sponge and then bending back the end of the wire stem sharply upon itself at a point just above where the hairs commence (Fig. 18 b).

By slightly curving the part of the stem carrying hairs, the brush will better adapt itself to the curved surfaces of porcelain dishes.

26. Test-tubes, Beakers, Funnels and Porcelain-dishes are washed in a stream of tap-water, their surfaces being rubbed meanwhile by the test-tube cleaner (Fig. 18 b).



Fig. 18.

TEST-TOBE CLEANER.

The brush must be cautiously moved as it approaches the bottom of a test-tube or boiling-tube, since the glass is very thin, and therefore easily broken by undue pressure.

Glass funnels which are used for qualitative analysis may hav

their stems cut off at about two inches from the shoulder (8), the sharp outer edge thus produced being at once removed by rubbing it with the face of a triangular file. The inside of the shortened fieck can then be easily cleaned by rubbing it with a strip of wood, or by inserting an ordinary tobacco-pipe cleaner from the shoulder of the funnel so as not to cut the hairs.

Occasionally the cleaning-brush fails to remove strongly adhering stains. Rubbing with the finger-end covered with a little moist-ened sea-sand or powdered pumice will usually be effectual in the case of an open vessel; and in the case of a bottle or flask a little water may be shaken round inside with sand; pumice powder, saw dust or small pieces of paper.

Or the stain may be removed by a suitable solvent, or by a liquid which renders the substance of the stain soluble by chemical change.

Hot dilute hydrochloric acid will remove most stains in this way'; but it is sometimes necessary to heat a little strong sulphuric or nitric acid in a vessel in order to cleanse it. Hot solution of caustic potash or of ammonia may also often be used with advantage to remove grease.

Apparatus should be washed as soon as possible after it has been used, since after the apparatus has stood aside for some time the surface is usually less easily cleansed.

Each article, after it has been carefully washed, and then rinsed out with tap-water, should be placed upside down in the small wicker-basket to drain. If it is required for immediate use for analysis and cannot be allowed to drain, it may be rinsed out with a little distilled water. This rapidly removes the tap-water which itself often contains dissolved impurities.

- 21. Draining-basket.—Before apparatus is placed in the locker, it should be made a rule to wash all glass and porcelain which is not in actual use, and then to place it in a wicker basket to drain. The basket is then put away with its contents. Dirty apparatus should never be kept in the basket.
- All iron apparatus should be carefully dried, and must be kept in a dry place to prevent it from rusting.

Metal apparatus must never be put into the basket.

22. Platinum Foil and Platinum Wire are cleansed by boiling them in hydrochloric acid, and rinsing off the acid with water. The wire should then be strongly heated for some time in the blow-pipe-flame, until, when it is wetted with pure strong hydrochloric acid and held in the Bunsen-flame, it no longer persistently colours the flame.

If the tip of the wire cannot be cleansed in this way, it should be cut off. Commercial platinum sometimes persistently colours the flame: wire made from such platinum is useless for flame.

coloration tests and for spectrum analysis.

Platinum-foil and wire should be kept in a small beaker containing strong hydrochloric acid, which has been previously diluted with sufficient water to prevent it from fuming. When the platinum is removed from the acid and has been rinsed with water, it will usually be found to be clean and to give no colour to the flame.

HEATING PORCELAIN AND GLASS.

23. A few general Precautions should be observed in heating glass and porcelain vessels, in order to guard against cracking them. The two following rules apply to both glass and porcelain:

A vessel containing a liquid must never be heated by the flame

above the level of the liquid inside.

A hot dry vessel must be allowed to cool before any liquid is poured into it, and before it is placed upon a cold surface.

24. Porcelain Vessels withstand higher temperatures than glass vessels, and are not so liable to be cracked by being suddenly heated or cooled.

*Porcelain Dishes are generally used for the purpose of evaporating liquids to dryness, or for concentrating them. They are supported on a pipe-clay triangle, which is placed upon a tripod-stand or a retort-stand. Porcelain dishes may be safely heated by a small naked flame.

Porcelain Crucibles are used for strongly heating solid bodies. Crucibles are supported in the same way as porcelain dishes. The flame should not be allowed to play steadily upon the bottom of the crucible so as to heat it suddenly, but the burner should be constantly moved slightly from side to side until the porcelain is hot.

The crucible should also be allowed to cool slowly on the triangle, since contact with a cold body is very apt to crack it. The hot crucible and cover are handled by the crucible-tongs.

25. Glass Vessels require to be Heated more cautiously than porcelain vessels. A large naked flame must never be allowed to play for any length of time on one part of the glass surface.

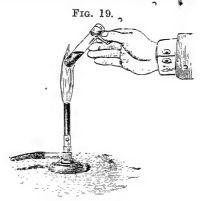
When a test-tube or boiling-tube is being heated, this local heating is prevented by holding the tube obliquely with the lower part in the flame, and either moving the tube slowly up and down, or shaking the contents of the tube (Fig. 19).

Test-tubes and Boiling-tubes.—Ordinary test-tubes are too narrow to contain safely a large quantity of boiling liquid, since the upper portion of the liquid is liable to be projected from the tube during ebullition. A small quantity of liquid may be boiled, and a larger quantity may be heated short of boiling, in a test-tube. But the broader "boiling-tubes" are to be preferred when a liquid is to be boiled.

Test-tubes, or boiling-tubes, which are not full of liquid, can

be held by the neck while they are being heated. They should be supported in an oblique position, so that the fingers are not over the flame (Fig. 19).

If steam or vapour is emitted in quantity, all risk of burning the fingers is avoided by bending round the neck of the tube a strip of folded glazed paper, or of leather, and pinching the ends of the strip together close to the tube, as is shown in the figure. This simple device serves better than a metal holder.



HOLDER FOR TEST-TUBE.

It is dangerous to use a strip of unglazed filter-paper, since if it becomes wetted it breaks and allows the tube to fall.

Crucible-tongs must never be used for holding test-tubes.

Glass Flasks, such as the wash-bottle, are most safely heated by placing them on a piece of wire-gauze on a tripod-stand over the Bunsen-flame (2), or they may be heated over the Fletcher flame (Fig. 2, page 3). If a large sand-bath, hot plate, or water-bath (Figs. 26, 28, pages 22, 23) is available, the flask may be heated on the hot surface or by the steam.

SECTION II

ANALYTICAL OPERATIONS, ILLUSTRATED BY **EXPERIMENTS**

29. Introductory Remarks .- Before the student tries the analytical reactions, he should become familiar with the operations which are commonly · employed in chemical analysis. He may obtain this knowledge by reading through the following descriptions, and then performing the illustrative experiments.

Each student should work independently.

In all analytical work the water used must be distilled water, and this

is conveniently kept in the wash-bottle.

A list of the chemicals required in this section will be found in par. 1408, and lists of the apparatus in pars. 1368, 1369.

DISTILLATION, DISTILLED WATER.

. 30. The Process of Distillation is often used to purify a liquid from solid substances which it contains in solution. The process consists in boiling the liquid and cooling or "condensing" the vapour. The solid substances are left behind in the vessel in which the liquid is boiled, and the condensed vapour forms the liquid "distillate."

The purification of spring water from the salts dissolved in it will serve as an example of this process.

31. Tests for Salts dissolved in Tap-water.—Certain salts, which are ordinarily present in spring-water, may be tested for as is directed below. But it must be understood that all the results which are described may not be obtained, since the soft water supplies of some of our large towns are almost free from dissolved solids. Rain water, or soft water, which gives no results with these tests, may be employed in qualitative analysis without previous distillation. A sample of water which contains saling impurities must be obtained for the following tests.

EXPERIMENT 1.—Make each of the following tests on a separate portion of the water contained in a test-tube. Chloride.—Add to some of the water several drops of diluta

nitric acid and of silver nitrate solution, and notice whether the water becomes turbid. Turbidity shows the presence of chloride in the water.

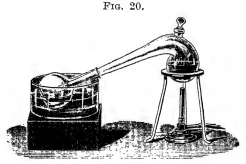
Calcium.—A separate portion of water in another test-tube may be found to become turbid on the addition of some solution of ammonia and of ammonium oxalate. Dissolved calcium salts produce this turbidity.

Sulphate.—The presence of sulphate would be indicated by a turbidity being produced on the addition to another portion of the water of a few drops of hydrochloric acid and of barium chloride solution.

Hardness.—Ascertain also whether precipitation of soap is caused by the water, owing to the presence in it of certain dissolved solid.

For this purpose dissolve a small shaving of soap by warming it in a little distilled water. Add a drop of this soap-solution to some of the water, which half fills a test-tube. Close the mouth of the test-tube with the thumb, and shake the water well.

If "hardening salts" are present no lather will form, but the water will be rendered more or less turbid by the separation of soap; but by the gradual addition of more soap-solution, the initial turbidity will be increased, and ultimately a lather will be produced when the water is shaken.



DISTILLATION OF WATER.

32. Distilled water and Tests for its Purity.—Distilled water is made in large quantities for use in various chemical processes. It must be substituted for ordinary water in the process of analysis, if the water-supply contains dissolved solids.

The student may use a glass retort and cooled flask (Fig. 20) to produce a little distilled water for the following tests, or he may use some of the distilled water supplied to the laboratory. An apparatus for preparing distilled water in larger quantity is described in paragraph 1377.

Q.A.

EXPERIMENT 2.—Distilled water, when it is examined by the tests described in paragraph 31, should remain perfectly clear Solution of ammonium exalate, of silver nitrate, of barium chloridand of ammonium sulphide when separately added to fresh portion of the water should produce no precipitate.

These tests prove the absence of calcium salts, of chloride, a sulphate, and of lead and iron salts respectively, any or all of which have been removed by distillation. Prove also that distilled wate at once gives a lather with soap-solution, and leaves no residue when it is evaporated to dryness (37).

SOLUTION.

dissolve in water, when they are stirred or shaken with that liquid. Salt and alum may be mentioned as examples. Certain other liquids may be employed instead of water; and if they cause solid substances which are immersed in them to become partially or entirely liquid, and to mingle uniformly with the liquid, they are said to "dissolve" the solids.

The liquid thus obtained is called a "solution" of the solid, and the liquid which dissolves the solid is termed the "solvent."

Further, a solid which dissolves in a liquid is said to be "soluble" in that liquid; if it does not dissolve, it is said to be "insoluble."

Thus when water is shaken with sodium chloride or common salt it dissolves the salt, yielding solution of sodium chloride; water is therefore called a solvent for sodium chloride, and this salt is said to be soluble in water.

The process of solution proceeds most rapidly when the solid substance is finely powdered before it is stirred in the liquid, since the largest possible surface of the substance is thus brought into contact, with the solvent.

The process is also much hastened by heating the solvent, partly because heat causes a rapid circulation of the liquid over the solid, and also because a solid substance is usually much more soluble in the hot liquid than in the cold.

Two kinds of solution may be distinguished, namely "simple solution" and "chemical solution."

34. Simple Solution occurs when a substance is dissolved by a liquid without undergoing permanent alteration in its composition.

A simple solution therefore generally possesses the taste, colour, and other general properties of the solid which it contains. It by "evaporation."

The solution of potassium nitrate or of copper sulphate in water is an example of a simple solution.

EXPERIMENT 3.—Place a small piece of potassium nitrate in a small clean beaker. Partly fill the beaker with water, and stir the solid about with a glass rod. The potassium nitrate will slowly dissolve in the water, and the solid will diminish in amount.

Now heat the liquid by placing the beaker on wire-gauze over a small flame; the solution will proceed much more rapidly.

Powder another piece of potassium nitrate by crushing it in a mortar, and then rubbing it round with the pestle. Place this powder in a beaker, pour in water, and heat the bottom of the beaker on wire-gauze by a small Bunsen-flame (Fig. 21 A, page 20). The potassium nitrate will dissolve much more rapidly than before; showing that the process of solution is accelerated by powdering the solid and by employing heat.

Mix these solutions, and preserve the liquid for future use.

EXPERIMENT 4.—Powder a little copper sulphate in a mortar, and transfer it to a small porcelain dish. Half fill the dish with water, support it upon a pipe-clay triangle placed on a tripod- or retort-stand, and heat it with a small flame. The blue copper sulphate will dissolve, yielding a blue solution. Keep this solution for future use.

These experiments (3, 4) are examples of simple solution. The first shows that a colourless solid gives a colourless solution, and the second that a coloured solid gives a coloured solution. This is generally true, and hence the presence of a coloured substance in a solution may be inferred if the liquid itself is coloured.

Moreover if a drop of the potassium nitrate solution is tasted, it will be found to possess the same taste as the solid.

Before a simple solution is analysed its colour is therefore usually noted, and its taste is occasionally tried with proper precautions.

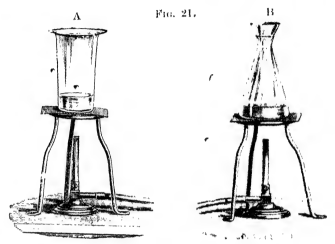
35. Chemical Solution differs from simple solution by producing a chemical change in the substance which is undergoing solution. The solution therefore contains a substance which differs in composition from the undissolved solid. Accordingly when the solvent is removed by evaporation the original substance is not obtained.

EXPERIMENT 5.—Place in a test-tube a small piece of marble or chalk, calcium carbonate, pour upon it a little water and heat the tube. The calcium carbonate will be found to be insoluble in water.

Now add to the water some hydrochloric acid: "effervescence," or escape of numerous small bubbles of gas, will occur. The piece

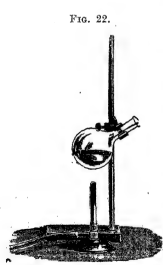
of calcium carbonate will meanwhile slowly diminish in size and will at last entirely disappear in the liquid, if sufficient acid is added.

EXPERIMENT 6.—Place in a test-tube a small piece of copper, and warm it with a little water: the copper will remain undis-



APPARATUS FOR DISSOLVING A SOLID.

solved. Now add to the water some nitric acid and heat: the copper will slowly dissolve, yielding a reddish-brown gas. The metal may be entirely dissolved, if sufficient nitric acid is employed.



DISSOLVING A SOLID,

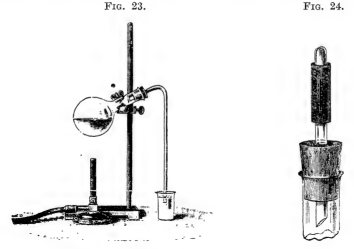
These are two examples of chemical The calcium carbonate is solution. changed by the hydrochloric into calcium chloride, and this substance, not the calcium carbonate, remains in solution. The is changed into copper nitrate, which is then dissolved by the water. solutions would therefore furnish on evaporation calcium chloride and copper nitrate respectively, and not marble or metallic copper.

It will be noticed that in each of these cases a gas is given off. This is a very usual, but not a universal effect during the process of chemical solution. The distinguishing fact is that the solid substance has undergone a change in composition in the act of passing into solution.

36. Apparatus for Solution of Solids.—As will have been already seen, the apparatus used for the solution of solids may be a test-tube, a porcelain dish, a beaker or a flask.

In Figs. 21 A, B, it is shown how loss by spirting from a boiling or effervescing liquid may be prevented by a clock-glass cover or a funnel. In Fig. 22 the inclination of the flask serves the same purpose, and in Fig. 23 the spirtings are collected in a small beaker containing water, and the water may be returned to the flask. These devices also tend to check the loss of the solvent by evaporation; they are, however, not usually necessary.

It is sometimes necessary to dissolve a substance without exposing it to contact with the air during the process. This may be effected by passing a stream of carbon dioxide through the vessel in which the solution is proceeding; or the air may be displaced from the flask by carbon dioxide, and



DISSOLVING A SOLID OUT OF CONTACT WITH AIR.

then a perforated cork may be at once fitted into the neck of the flask. The perforation carries either a bent tube with its end just immersed in a small quantity of distilled water (Fig. 23) forming a water valve, or a short tube surmounted by a piece of rubber tube which is closed with a solid glass stopper: the rubber tube has a slit cut in it by a sharp knife (Fig. 24), and this slit gives egress to any gas or vapour from within, but excludes the entrance of air from without.

EVAPORATION AND CRYSTALLISATION.

37. The Process of Evaporation.—When it is necessary to separate a solid substance from the liquid in which it is dissolved, the liquid is boiled away as vapour or "evaporated." The solid

substance is then left behind in the vessel. The liquid is usually removed by heating the solution in an open porcelain evaporating dish over the Bunsen-flame.





EVAPORATION IN DISH.

EXPERIMENT 7. Poul the potassium nitrate solution (Exp. 3, page 19) into a porcelain evaporating basin, and heat it over the Bunsen flame until the water has been nearly removed:

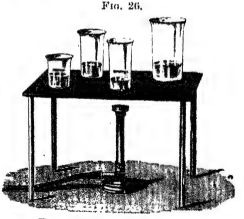
Then make the flame smaller, and continue heating until the water has disappeared. Solid potassium nitrate will be left in the dish.

Towards the end of the process of evaporating a solution, a small flame should always be used; and in order to prevent the substance from spirting out of the dish, the flame is moved about, or the dish may be covered with a funnel (Fig. 25) or with a circular filter-paper.

The process of evaporation in a shallow dish over a naked flame is shown in Fig. 42, and the covering of the liquid during the latter stage of the process is indicated. Fig. 43 illustrates another method of evaporating on a freated iron plate, the liquid being contained in a beaker instead of in a dish; the nearer the beaker is placed to the part of the plate which is over the flame, the more rapid will be the evaporation.

38. Evaporation at Steam-heat. — It is often necessary to evaporate more slowly at steam-heat, in order to prevent the spiriting of the liquid or the overheating of the solid. For this purpose the dish is placed over boiling water upon a waterbath, steam being in this case the heating agent.

A simple form of water-bath is shown in



EVAPORATION IN BEARERS.

Fig. 27. It consists of a copper vessel which is rather more than half-filled with water, and is heated by a Bunsen-burner. The

upper portion is fitted with flat sheet-copper rings of gradually diminishing diameters, and the top of the bath can thus be made

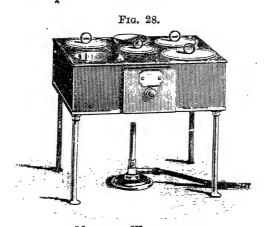
to support vessels of various sizes. A more complex form of copper water-bath is shown in Fig. 28. In the upper surface of this bath there are holes of various sizes, designed to fit vessels of different dimensions. When these holes are not in use, they are covered by lids as is shown in the figure. This form of waterbath is usually fitted with the automatic water supply represented in Fig. 41 (page 33).

A simple water-bath may be improvised from a beaker partly filled with water and heated by a Bunsen-flame (Fig. 29, page 24).



SINGLE WATER-BATH.

A few small pieces of paper may be thrown into the water in order to prevent it from bumping during the boiling.



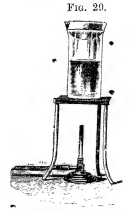
MULTIPLE WATER-BATH.

When the evaporating dish is placed on the top of the beaker the escape of the steam may possibly be prevented. If this should be the case, some strips of paper may be inserted between the bottom of the dish and the edge of the beaker.

In a large laboratory used by many students the steam-baths are often provided by openings closed with lids (Fig. 45) upon the top of a range of steam-ovens in the combined arrangement shown in Fig. 75 (1377).

39. Concentration and Crystallisation. Sometimes only a part of the liquid is evaporated for the purpose of "concentrating" the solution.

When a hot solution has been sufficiently concentrated, it will



SIMPLE WATER-BATH.

frequently deposit "crystals" of the dissolved substance as it cools.

Soluble substances are often dissolved, and then crystallised from their solution in this way, in order to free them from impurities.

EXPERIMENT 8.— Concentrate the copper sulphate solution which was prepared in Exp. 4 (page 19), and allow the liquid to cool. If sufficient water has been evaporated, crystals will form.

In order to ascertain whether the hot solution will crystallise on cooling, pour about two cubic centimetres (cc.) of the solution into a test-tube and cool the test-tube in water; if crystals

form, the rest of the solution may be allowed to cool in the air.

Keep the copper sulphate solution in the dish for future use.

PRECIPITATION.

40. Precipitation by a Liquid.—When two clear and transparent solutions are mixed together, they frequently become more or less turbid or opaque, owing to particles of a solid substance being formed in the liquid. A solid substance thus produced is called a "precipitate," and the process of producing a precipitate is termed "precipitation."

A substance is frequently removed from solution by causing it to form an insoluble compound or precipitate. The substances which are added for this purpose are termed "reagents" or "precipitants": they are commonly liquids.

In producing a precipitate, care must be taken that the two solutions are well mixed. Mixture may be effected by warming the bottom of the test-tube in the flame, by stirring the liquid with a glass rod, or by pouring the liquid from one vessel to another.

A precipitate often appears more rapidly when the liquid is vigorously shaken, or when it is warmed.

EXPERIMENT 9.—Pour some barium chloride solution into a test-tube, and add ammonium carbonate solution. A white precipitate of barium carbonate will form. Keep the precipitate and liquid in the test-tube for future use.

In this instance barium chloride and ammonium carbonate may be readily dissolved in separate portions of water. But, if these solutions are mixed, two different substances, ammonium chloride and barium carbonate, are produced. The former of these remains dissolved in the water; but the latter is insoluble, and therefore separates as a fine powder in the liquid.

Precipitates differ much in colour and in general appearance and properties; they are therefore frequently produced, by the addition of suitable reagents, in order to show the presence of a substance. Precipitates are also produced for the purpose of

separating one substance from another.

The appearance of a precipitate is usually described by its colour and by its condition. It is "flocculent," if it forms in flock-like masses; "crystalline," if it consists of small particles which are seen to be crystals under a lens or microscope; "gelatinous," if it is jelly-like in its consistency. The formation of a slight precipitate may cause only a "turbidity" in a liquid.

The day-light colour of a precipitate is always given in the following text, and this may be changed when the precipitate is seen in some kinds of artificial light. The true colour, however, is seen at night if the precipitate is illuminated by the electric arc, or by a piece of burning magnesium ribbon.

41. Precipitation by a Solid Reagent is occasionally resorted to. Thus a metal is not unfrequently precipitated from the solution of its salt by the immersion in it of another metal.

EXPERIMENT 10.—Dip a clean penknife-blade into some of the copper sulphate solution (Exp. 8), to which a few drops of sulphuric acid have been added. After a short time metallic copper will be precipitated from the solution and will cover the iron as a red film.

DECANTATION AND FILTRATION.

The Processes of Decantation and Filtration serve to separate a precipitate from a liquid in which it is suspended. Decantation is sometimes preferred because with suitable precipitates it is more rapidly and simply carried out than filtration, and it leaves the precipitate in the vessel ready for immediate further treatment.

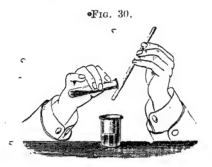
42. The Process of Decantation serves to separate a heavy precipitate, which settles rapidly, from the liquid out of which it has subsided.

For this purpose, the vessel containing the liquid and precipitate is allowed to stand until the precipitate has settled. The liquid

is then carefully poured off, or "decanted," by gently inclining the vessel. A wetted glass rod, pressed against the edge or lip of the vessel (Fig. 30), helps to prevent the precipitate from being disturbed while the liquid is flowing away from it.

If the process of decantation is carefully performed, it will separate the liquid almost completely and in a clear condition.

EXPERIMENT II.—Add boiling dilute sulphuric acid to some



DECANTATION OF A LIQUID.

boiling solution of barium chloride. A heavy precipitate of barium sulphate will be formed. This precipitate may be readily separated from the liquid by decantation. Keep the precipitate in the test-tube for future use.

43. Filtration.—The liquid containing the precipitate is poured into a cone of porous paper. The liquid itself runs through the pores of the

paper, but the solid particles of the precipitate are retained upon the surface of the paper.

The paper employed is called "filter-paper," and the liquid which has passed through the filter is termed the "filtrate."

A filtrate may frequently be coloured by some substance dissolved in it, but it should be perfectly free from turbidity caused by solid particles suspended in it.

The ordinary process of filtration employed in chemical analysis is described below, methods of filtration on a larger scale are described in paragraphs 1391, 1392.

Prepare a filter for filtration as is directed below.

EXPERIMENT 12.—Measure a glass funnel (Fig. 48) along its sloping side from shoulder to rim. Select a circular filter paper, the radius of which is somewhat less than this in length: fold it across into a semicircle, then fold it again at right angles into a quarter of a circle; now open out either of the sides, so as to form a little conical bag (d, Fig. 32). This forms an ordinary filter.

Filters 3½ and 4½ inches (9 and 11 cm.) in diameter are convenient for ordinary purposes

meter are convenient for ordinary purposes.

If ready-cut filters of suitable size are not at hand, cut from a sheet of filter-paper a smare piece, the edge of which is rather less than double the





FUNNEL.

length of the side of the funnel. Fold it over along the line (a, Fig. 32, then again along the dotted line (b). This gives a square (c) which at one angle has four free corners; these are removed by cutting with a pair of scissors along the curved dotted lines shown in (c).

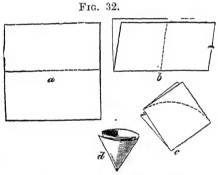
The filter is now made and only requires to be opened. By separating the curved edges so that they form a circle, three remaining on one side of the circle and one on the other, a little closed pointed paper bag is formed (d).

A filter of the same shape, but of uniform thickness, may be made from half an ordinary circular filter-paper, or from half of the circular flat paper formed by opening out the filter (d). This semicircular paper is folded into a quadrant shape, and the two radial edges are doubled over together several times, the fold being pressed down with the finger nail. The unused half of the filter (Fig. 32, d) is absent in the filter which is formed by opening this paper, and filtration is more rapid in consequence.

The folded filter is now gently pressed with dry fingers into the

dry funnel, and the folding is altered if necessary until the paper fits the glass closely. It is then moistened all over with distilled water from the wash-bottle and is ready for ase.

This preliminary moistening of the filter-paper must not be neglected, since if the liquid which contains the precipitate is poured upon a dry filter, some particles of the pre-



FOLDING AND CUTTING FILTER.

cipitate may lodge between the fibres of the paper; the paper then shrinks on being wetted inclosing the solid particles, and these choke the pores of the filter and seriously retard the filtration.

During the filtration of a liquid the funnel may be placed with its neck in a test-tube, which is supported in the test-tube stand. Care must, however, be taken that the inside of the upper part of the test-tube is dry, and that there is a space between the neck of the funnel and the inside of the tube; since, if the egress of air from the test-tube is prevented, the filtration will be stopped.

The funnel is therefore preferably supported upon the ring of a wooden filter stand (Fig. 33, page 28); and the filtrate is received in a small beaker standing beneath the funnel, instead of in a test-tube.

As has been already stated, the upper edge of the filter must always be below the rim of the furnel; and when the liquid, which is to be filtered, is being poured into the filter, care must be taken not to let it reach the upper edge of the filter-paper.

The precipitate should not more than half fill the filter, else it cannot conveniently be washed.

EXPERIMENT 13.—Pour the liquid containing the precipitate of_ barium carbonate, from Exp. 9 (page 24), through the fifter. If the above directions have been carried out, and no hole has been made in the paper, while it is being prepared or fitted into the funnel, the filtrate will run through perfectly clear, leaving the barium car-

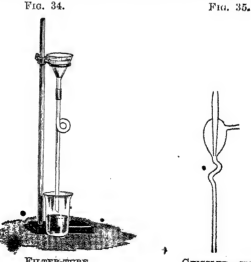
FILTER-STAND.

bonate on the filter. Keep the filter and precipitate in the funnel for future use.

Sometimes the filtrate passes away from the filter in a turbid condition.. In order to remove this turbidity, the turbid portion of the filtrate must be poured once or twice through the same filter. Or the turbidity may be prevented by pouring the liquid at once through a double filter, which is prepared by folding two filter-papers together: as one: a double filtration is thus secured in one operation.

It should be borne in mind that a hot liquid passes through the filter more quickly than a cold one. Hence, if it is not inadmissible for other reasons, a solution should always be heated before it is filtered.

44. Accelerated Filtration.-Methods for expediting the filtration of arge quantities of solids from liquids are described in pars. 1391, 1392. But



FILTER-TUBE.

GEISSLER-ASPIRATOR

it is often desirable to hasten the process of filtration of a liquid in an ordinary funnel, more especially when a floceulent precipitate is being separated from the liquid. This may be effected either by attaching a suitable tube to the funnel (1), or by means of a filter-pump or an aspirator (2).

(1) One of the simplest means of increasing the rate of filtration is to attach to the funnel a glass tube, about 2 feet (69 cm.) long, bent as is shown in Fig. 34. As soon as the filtrate has filled this tube, the weight of the column of liquid tends to draw the liquid through the filter, and the rate of filtration is accordingly increased.

Another simple method is to attach to the funnel a straight piece of glass about 2 feet (60 cm.) in length and 1 millimetre in bore. The weight of the liquid column which is suspended in this tube pulls the liquid through the filter. \bullet

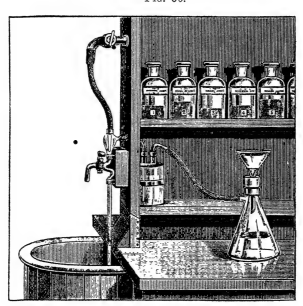


Fig. 36.

WATER FILTER-PUMP, OR ASPIRATOR.

(2) A Filter-pump, or Aspirator, which applies suction to the neck of the funnel, is frequently used to hasten filtration.

The accompanying illustration (Fig. 36) shows a convenient arrangement for this purpose. The glass Geissler-aspirator (Fig. 35) is shown over the sink, connected by stout rubber tubing with the high pressure water-tap. When the water is turned on and flows through the aspirator, air is sucked from the thick-glass conical tubulated filter-flask; and the atmospheric pressure, acting on the liquid in the funnel, is thus utilised for forcing the liquid through the filter.

The two-necked Woulffe's bottle is interposed between the aspirator and the filter-flask in order to prevent the water in the aspirator from entering the flask.

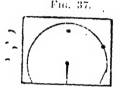
The neck of the funnel is fitted by a rubber cork into the neck of the conical filter-flask; and in order that the pressure which is exerted on the point

of the filter-paper may not burst it, either toughened filter paper may be used, or the ordinary filter-paper may be supported beneath by a small cone of parchment-paper or of platinum.

The cone is cut from a piece of platinum foil or parehment paper measuring about 11 - 1f inch (Fig. 37). A hole is pierced by a pin near the middle. a cut is made from the edge to this hole, and the corners are removed by cutting round in a circle which has the pin hole for its centre. If the metal

foil is used, the spring is then removed by heating it to

reduess and cooling it slowly.



PLATINUM FILTER-CONE.

A cone (Fig. 32, d, page 27) is now made by folding over, and is fitted exactly into the hortom of the fulinel. A filter-paper is then folded as usual and is titted closely into the funnel, both filter and funnel being dry.

Before the filter is used for filtration it is meistened with water, and is then gently pressed, where necessary, with the fingers until it adheres to the funnel; the upper edge of the filter especially requires to be carefully pressed against the glass in this way. The filter is now tested,

by filling it with distilled water and setting the pump in action.

If the paper fits properly and has been pressed into contact with the funnel, no air-bubbles will be sucked down the neck of the funnel until the whole. of the water has passed through the filter. Any air leakage noist be stopped By pressing the paper against the funnel where such leakage occurs,

Besides expediting the filtration and washing of precipitates, such an arrangement also enables the precipitate to be dried to a great extent by sucking air through it after the liquid has passed away.

The aspirator may be replaced by other forms of apparatus for producing suction. A common hand-syringe may be used, but this is liable to be attacked by acid vapours if it is made of brass.

A Special form of Receiver is represented in Fig. 38 to serve as a substitute for the conical flask. It consists of an ordinary separating funnel, F. which is closed with a doubly perforated rubber cork.

One perforation carries the funnel, which is used for ultration. Into the other is inserted a branched tube, one limb of which (h) is connected with the filter-pump, while the other (a) can be opened to the atmosphere by turning the stopcock.

Fig. 38.

SPECIAL RECEIVER FOR THE FILTRATE.

The use of this receiver renders it possible to withdraw a portion of the filtrate during the process of filtration, by opening simultaneously the stopeocks a, c and stopping the action of the filterpump.

WASHING THE PRECIPITATE.

45. The Washing of a Precipitate, after it has been separated from the liquid in which it was suspended, is usually necessary in order to free the precipitate from the adhering solution.

If the precipitate has been filtered, it is washed upon the filter (46); if it has been separated by decantation, it is usually washed by decantation •(47).

*46. Washing the Precipitate on the Filter.—The following directions will serve to explain how this process is carried out, and will indicate what precautions should be observed in order to secure success. The washing process may be hastened by employing the devices which have been already described in par. 44.

EXPERIMENT 14.—Support the funnel, which contains the fifter and the barium carbonate precipitate from Exp. 13 (page 28), in a filter-stand above a beaker or flask (Fig. 33, page 28). Blow into the filter a fine stream of hot distilled water from the wash-bottle (17), so directing the jet as to stir up the precipitate. Fill the paper in this way with water to within a short distance from its edge, and let the water run through perfectly, and repeat these processes three or four times, letting the water run through completely each time before adding a fresh quantity.

The precipitate and the filter should now be free from everything soluble in water, and the water which passes through the filter will therefore be tasteless.

Ascertain whether the washing is complete, by collecting the last few drops of the washing-water in a clean test-tube, and testing whether it contains any chloride by adding to the liquid a drop of silver nitrate solution; no turbidity must be produced. If turbidity appears, the washing must be continued and the washingwater must be tested again.

The process of washing must be continued, until silver nitrate produces no turbidity when it is added to the last portions of the water running from the funnel.

Reserve this precipitate in the filter for further use.

47. Washing the Precipitate by Decantation.—This method may be adopted with a suitable precipitate, and may often usefully precede the separation of the precipitate from the liquid by filtration, the earlier stages of washing being carried out by decantation.

EXPERIMENT 15.—The precipitate of barium sulphate from Exp. 11 (page 26) may be washed by decantation as follows.

Hot water is poured into the vessel containing the precipitate, and the water is shaken, stirred, or boiled with the precipitate, which is then allowed to settle. As soon as the water has become clear, it is poured off as completely as possible in the way shown in Fig. 30 (page 26).

By repeating this washing process several times; with the addition of fresh portions of boiling distilled water, the precipitate may be entirely freed from the adhering solution.

While the washing-water is being poured off it should be occasionally tested with blue litmus paper, in order to ascertain when it ceases to redden the paper and is therefore free from acid. As soon as the washing-water is quite free from acid, the precipitate may be considered to be thoroughly washed.

Reserve this precipitate for further use (56), after filtering it

off on a small filter, and putting it aside to dry.

It is usual to Ascertain when the Washing of a Precipitate is Complete, by testing the washing-water for some soluble substance which is being removed by the washing. That substance is usually tested for which is most readily detected. Experiments 14 and 15 serve to illustrate this statement.

DRYING THE PRECIBITATE.

48. A Precipitate may be dried by supporting the funnel on a hollow tin cone or cylinder, called a "filter-dryer" (Fig. 39)

Pig. 39.

FIGTER DRYER.

which is placed on a piece of iron wire gauze on a tripod-stand, and heated over the flame of a rose-burner turned very low; or the filter dryer may be placed upon a heated iron plate or sandbath.

The filter may be dried more quickly by allowing it to drain for a time, and then opening it out and spreading it upon a piece of wire-gauze supported upon a tripod-stand or upon the ring of a retort stand, a small flame from the rose-burner being placed at some distance beneath it. The hand ling of the moist filter is facilitated if it has first been laid upon a pile of several filters, which will absorb

the water. The filter is by these means exposed to a current of hot air, which

rapidly dries the precipitate. Care must be taken to regulate the heat so as not to char the filter.

The Steam-oven may be used to secure a more gradual but safer process of drying. The funnel is placed in the steam-oven as is shown in Fig. 40, but the conical support is usually unnecessary, since most ovens have perforated shelves and the funnel is supported by passing its stem through one of these perforations.

A combined system of steam-ovens and water-still is shown in Fig. 75 (1377) for use in a large laboratory, and in some laboratories the steamovens are heated by steam supplied from a large general boiler. These steamovens will be available for drying precipitates.

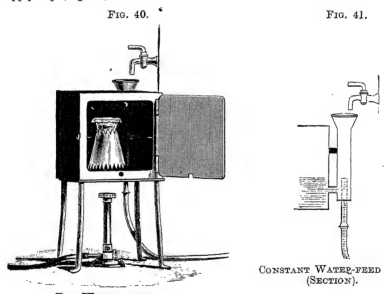
But if this general system is not available the separate steam-oven shown

in Fig. 40 may be used. In this oven water is boiled in the space between the double-cased copper oven, the escaping steam being condensed and returned to the water-jacket by connecting the steam-outlet with a long piece of "composition" tubing fixed on the laboratory wall, and thus maintaining the water level. This tube must be sufficiently long to condense all the steam generated when the water is gently and steadily boiling.

Another method of maintaining the water-level is shown in section in Fig. 41, where a slow drip from the water-supply pipes falls into a funnel feed-tube at the side of the oven, any excess of water escaping by a central

out-flow pipe into the waste.

It is well to avoid a constant feed of hard water to the steam-oven, since the incrustation which gradually forms in the water-space is removed with difficulty. Accordingly if the oven is not heated by steam, or the interspace constantly fed, as directed above, by the return of the condensed steam, a raised vessel containing soft or distilled water may be used to feed the supply tap (Fig. 41).



THE WATER-OVEN.

The air-inlet pipe to the bottom of the oven passes through the hot water and delivers hot air in the interior, an outlet for air being provided at the top of the oven. It is convenient to have a piece of glass let into a metal frame for the door, so as to render the interior visible without opening the door.

The Air-oven (Fig. 42, p. 34) may be employed instead of the steam-oven. It is similar in construction, but is heated directly by a gas-flame placed beneath it, the temperature of the interior being suitably adjusted by regulating the flame.

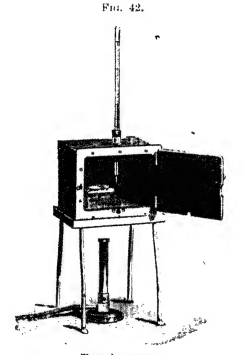
The requisite temperature may be maintained by occasional observation of the thermometer, or a special thermal gas regulator may be enclosed in the interior of the oven, and will serve to automatically control the gas-supply.

The air-oven can be used when drying temperatures above 100° C, are required; but it must be remembered that the metal sides of the bath are much hotter than the internal air, and that the drying substance should therefore be supported out of contact with the oven, as is shown in the figure, if it is to be heated only to the temperature registered by the thermometer.

The air-oven may be heated electrically instead of using a gas-burner, and with suitable resistances a considerable range of temperature is thus obtainable.

REMOVING THE PRECIPITATE FROM THE FILTER.

49. Removal of Part of the Precipitate from the Fifter. A small portion of a moist precipitate may be taken from the filter by dipping the end of a glass rod into it. If a watch-glass, or the



THE AIR-OVEN.

interior of a test-tube, is then touched with the end of the rod, the precipitate is deposited upon the surface of the glass, and may be subjected to further test or examination.

A larger quantity of the precipitate may be taken out of the filter by means of a glass spatula, made by flattening out the end of a glass rod which has been softened in the flame.

50. Removal of the Whole of the Precipitate from the Filter.—If the precipitate is to be removed from the filter as com-

pletely as possible, several methods are available. One or other of those described below under a, b, c, d, e, and f, must be chosen according to circumstances. Of these the methods described under a, e and f are in most common use.

- (a.) Washing down the Precipitate through the neck of the Funnel.—A hole may be made in the bottom of the filter by means of a thin glass rod (15), which is pushed down through the neck of the funnel. The precipitate is then easily washed down into a vessel, placed beneath the funnel, by a fine stream of liquid from the wash-bottle.
- (b.) Washing out the Precipitate from the Rim of the Funnel.—The funnel may be held with its neck horizontal, and

Fig. 43.



WASHING THE PRE-CIPITATE OUT OF THE FILTER.

with its rim just inside the edge of a porcelain dish (Fig. 43). The precipitate is then washed out of the filter by directing a fine stream of water from a wash-bottle against the inside.

(c.) Removal of the Filter with the Precipitate from the Funnel, followed by rinsing off the Precipitate.—The filter with the precipitate is allowed to remain in the funnel for some time, in order to permit as much water

as possible to drain away. The filter is then carefully taken out of the funnel and is further dried, if necessary, by laying it upon several folds of filter-paper. After the portions which contain no precipitate have been removed, the filter is spread out inside a porcelain dish.

The liquid, with which the precipitate is to be treated, is then poured into the dish; and this liquid is shaken round in the dish, and the filter is carefully rubbed with the rounded end of a glass rod, until the precipitate has been removed from the paper. With a little care this may be effected without tearing the paper. The filter-paper is then removed from the liquid by means of the glass rod.

(d.) Removal of the Filter with the Precipitate, followed by Scraping off the Precipitate.—If it is undesirable to add a liquid to the precipitate upon the filter, the precipitate is allowed to drain for a short time; it is then further drained, if necessary, by removing the filter from the funnel and laying it upon several filter-papers.

The filter is then spread out upon a flat piece of glass, and the precipitate is carefully scraped off with a glass rod which is pressed

flat upon the paper, or with a small glass spatula (49).

This method is usually the most imperfect, but is frequently the best for other reasons.

(e.) Removal of the Precipitate by Dissolving it in the Filter.—If a precipitate is to be dissolved off the filter, the liquid, which is to be used as a solvent, is heated and is then poured upon the precipitate. The solvent will run through the filter into a vessel placed below the funnel, and will take with it the precipitate in solution.

After the Equid has passed through the filter, it should be heated again and once more poured upon the precipitate, if the latter is not entirely dissolved. The liquid is reheated and returned to the filter in this way as long as anything remains undissolved. If any portion of the precipitate still remains, it must be removed by using a fresh portion of the solvent.

EXPERIMENT 16. Remove in this way the precipitate of barium carbonate (Exp. 14, page 31) from the filter, by means of hot dilute hydrochloric acid.

(f.) Removal of the Precipitate by Rinsing it through the Filter with the Solvent.—A precipitate may also be removed from the filter by means of the liquid with which it is to be treated or dissolved, in the following manner.

The funnel is supported over a suitable vessel. The liquid is poured in and is quickly stirred up with the precipitate by means of a thin glass rod. The bottom of the filter is then pushed out through the neck of the funnel by the glass rod. The liquid will now flow through the neck of the funnel, carrying the precipitate with it.

If some of the precipitate remains on the filter, the liquid must be poured again through the funnel, and by repeating this process all the precipitate may be removed.

USE OF TEST-PAPERS.

51. Litmus and Turmeric Test-papers.—The two vegetable colouring substances litmus and turmeric undergo marked changes of colour when they are acted upon by certain substances. These changes serve to indicate whether a liquid or gas is "acid," or "alkaline": if no change occurs, it may be inferred to be "neutral."

The analyst is usually provided with slips of paper which have been stained with solutions of these colouring substances. These slips of paper are known as "test-papers". The different papers are easily distinguished by their colours. The following experiments will explain their use. 52. Acid, Alkaline, or Neutral Reaction in a liquid is ascertained by the behaviour of the liquid with these test-papers.

EXPERIMENT 17.—Dip a clean dry glass rod successively into dilute hydrochloric acid, into ammonium hydroxide solution, and into barium chloride solution. After each immersion of the rod, draw its end, wetted with the solution, successively across blue litmus-paper, red litmus-paper, and yellow turmeric-paper. The following results will be obtained.

Hydrochloric acid will redden blue litmus, but will not alter the colour of red litmus or of turmeric; it possesses an "acid reaction."

Ammonium hydroxide solution will not affect the colour of blue litmus, but will turn red litmus blue and turmeric brown; it shows "alkaline reaction."

Barium chloride solution will not affect the colour of any one of the test-papers: it is "neutral."

· Each of these liquids is typical of a large class of substances, in so far as its behaviour with these test-papers is concerned.

53. Neutralisation: Addition of Acid or of Alkali in Excess.—It is occasionally necessary to exactly neutralise an acid liquid by means of an alkaline liquid or vice versâ. The acid or alkali must also often be added in excess. The following experiment explains how these processes are effected.

EXPERIMENT 18.—Pour some dilute hydrochloric acid into a porcelain dish. Add solution of ammonium hydroxide drop by drop, constantly stirring the liquid with a glass rod and touching the wet end of the rod against a piece of blue litmus-paper.

By proceeding cautiously, a point may be reached at which the liquid does not affect the colour of either blue or red litmuspaper; the acid is then said to have been "neutralised" by the alkaline liquid.

The latter part of the process of neutralising the acid liquid should be completed by the addition of the alkaline solution in a very dilute condition, since the addition of an excess of the alkaline solution is thus more readily avoided.

Add another drop of ammonium hydroxide: the liquid will now have acquired an alkaline reaction: the alkaline solution is accordingly said to have been added "in excess."

By a similar procedure an alkaline solution may be neutralised with an acid, and may then be mixed with acid in excess or acidified.

54. General Remarks on Test-papers.—In choosing between red litmus and turmeric test-papers for the detection of alkaline reaction, it should be remembered that turmeric-paper is usually

more sensitive than red litmus paper, since red litmus paper may contain an excess of acid.

A turbid liquid will often form a deposit when it is placed upon the test-paper, and this may conceal the colour of the paper. The colour will, however, usually be visible on the back of the paper if it is unglazed. If the paper is glazed, the colour will be seen after the deposit has been washed off by a stream of distilled water from the wash bottle.

Test papers must always be kept in a stoppered bottle, in order to guard them against undergoing change by the action of gases contained in the laboratory air. The bottle should be shielded from light as much as possible, since these vegetable colours fade taider the action of sunlight or strong daylight: this is especially the case with turmeric paper.

SIMPLE TAXITION

55. Solid Substances are Heated strongly, or Ignited, in order to study the effect of heat upon them, since by this means their composition or nature is frequently revealed more or less completely. Ignition also serves to remove volatile from non-volatile substances.

It is a necessary condition of the process of "simple ignition" that the substance, which is being heated, shall not suffer change by contact either with the flame, or with the substance of the vessel or support, or by admixture with chemical reagents. It is usually, however, subjected to change by contact with the oxygen of the air.

EXPERIMENT 19. Heat some white zinc oxide in a small ignition-tube (13). The hot substance will be lemon yellow, but the colour will change to white again as the oxide cools. These changes of colour are characteristic of zinc oxide.

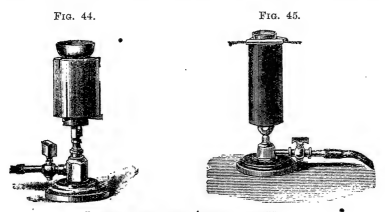
In the early stage of the heating, drops of water will probably condense in the upper part of the tube: this shows that adherent moisture was present with the oxide. Ignition in a tube is often employed to detect the presence of water in a substance.

EXPERIMENT 20.—Heat a little mercuric oxide in a small testtube or ignition-tube. After a short time minute shining drops of the metal mercury will be seen in the upper part of the tube. If a splinter of wood with a spark at its end is now held in the tube, it will burst into flame. The behaviour of this red powder when it is ignited, shows that it contains the elements mercury and oxygen. EXPERIMENT 21.—Heat a few particles of charcoal powder on platinum foil, by directing the tip of the blowpipe-flame against the under surface of the foil. The charcoal will be slowly burnt away by the exygen of the air, showing that the black substance consists of carbon.

EXPERIMENT 22.—Heat a small piece of ammonium chloride in an ignition-tube. The ammonium chloride will pass off as vapour or "volatilise," and the vapour will cool in the upper part of the tube forming a coating or "sublimate." This proves that the salt is volatile.

EXPERIMENT 23.—Heat in the same way a small piece of **potassium chloride**. This will melt or "fuse," but will not pass off as vapour. It will become solid again on cooling. Potassium chloride is almost non-volatile at the flame temperature.

Hence if a mixture of ammonium chloride and potassium chloride is heated on platinum foil, the potassium chloride alone will be



IGNITION OVER THE ARGAND-BURNER,

left in the residue, since the ammonium chloride will be removed as vapour.

If the quantity of the substance to be ignited is large, it is usually heated in a porcelain or platinum vessel over the Bunsen-, Teclu-, or blowpipe-flame (Fig. 64, 1308).

Small quantities of substance may be conveniently ignited in little porcelain or platinum capsules over an Argand burner from which the gauze top has been removed (Figs. 44, 45).

The luminous flame must not be allowed to touch the capsule, since it soils the surface and also damages platinum.

TREATMENT WITH FUSED REAGENTS.

56. A-Substance is often Heated with certain Solid Reagents until they fuse or melt, in order to cause chemical changes to occur between the substance and the reagents.

EXPERIMENT 24.—Mix a little of the dry barium sulphate (Exp. 15, page 31), which is insoluble in water and in acids, with a mixture of sodium and potassium carbonates, and heat the mixture on platinum foil (see *Note* below) until the alkaline carbonates fuse. Maintain the salts in a fused condition for several minutes. The following change will then have taken place:—

$$BaSO_4 + Na_2CO_3 = BaCO_3 + Na_2SO_4$$
.

 σ Now treat the cool product with water. The sodium sulphate will be dissolved. Separate the residual barium carbonate by filtration, and wash it. Then treat it with dilute HCl: the insoluble carbonate will dissolve as chloride: BaCO₃ + 2HCl BaCl₂ + H₂O + CO₂. The original barium sulphate is thus converted into chloride which is obtained in solution.

A substance which cannot be dissolved in water or in acids may frequently be converted into a soluble substance in a similar way to that just described.

The mixture of alkaline carbonates, which is used for the purpose, is generally known as "fusion mixture."

Note.—In this experiment, and in other experiments for which platinum foil is used, the platinum may be replaced by a small porcelain crueible or by a fragment from a broken porcelain dish. If porcelain, however, replaces platinum, a somewhat more powerful burner is necessary to produce the fusion.

Sometimes the treatment with fused reagents yields a characteristic result, which serves to detect the presence of a substance.

EXPERIMENT 25.—Place a little powdered sodium carbonate and potassium nitrate on a piece of platinum-foil (see Note above), and then add a minute quantity of manganese dioxide. Hold the foil with crucible-tongs in the top of the Bunsen-flame, or heat the under surface of the foil with the point of the blowpipe-flame. The mass will soon melt or fuse. Keep the substances in a melted state for a short time, and then allow the mixture to-cool.

A bluish-green mass will remain upon the foil, the colour of which is due to the sodium manganate which has been formed by the fusion. This colour is produced only when manganese is present, and the above process therefore serves as a test for that metal.

In the above experiments (19 to 25) the gases of the flame are

not allowed to take any part in the change which is described. In the following cases the flame-gases aid in producing the reaction.

• FLAME REACTIONS WITH REAGENTS.

57. The Gases of the Blowpipe-flame (II) or of the Bunsen-flame may play the part of Reagents in affecting the composition of the substance which is being heated. In order to understand the nature of these changes the structure of the flame must be carefully observed.

In a well-formed blowpipe-flame or Bunsen-flame two parts may be distinguished: the "inner-flame," which is blue and pointed; and around and beyond this an almost colourless flame, which is commonly known as the "outer-flame."

For ordinary mouth-blowpipe operations a luminous gas flame should be used, and the size of the flame should be so adjusted that both the inner and outer blowpipe flames are sharply defined; or the Bunsen-flame may be used after closing the air-holes more or less completely.

The Inner-flame is often called the "reducing or deoxidising flame," because it separates or reduces metals from their oxides and from many of their compounds, and in general deoxidises substances. This deoxidising power is due to the action of the hot but incompletely burnt combustible matter, which is present in this part of the flame.

The inner flame is obtained of large size by placing the tip of the blowpipe nozzle only in the outer edge of the flame.

The Outer-flame is also known as the "oxidising-flame," because it changes metals into their oxides, and oxidises many other substances, by supplying them with oxygen at a high temperature.

The size of the outer-flame is increased by pushing the nozzle of the blowpipe into the interior of the Bunsen-flame.

In the following processes the results obtained will usually vary when the substance is heated in the inner-flame or in the outerflame.

• 58. The Borax-bead—Fusion with Borax.—When certain metals and their compounds are fused with borax, they are converted into coloured borates. The colour of each of these borates is characteristic, either in itself or in the changes which it undergoes in the outer and inner flames.

The method of making the borax-head is explained in Exp. 26. Its use for testing is illustrated by Exp. 27.

EXPERIMENT 26. Place a little powdered borax on a watchglass. Moisten the loop of a piece of mounted platinum-wire (14). and dip it into the borax. Then hold the loop with the adhering powder in the outer blowpipe-flame until the borax melts.

If the bead thus formed does not fill the loop, dip the melted bead again into the borax; more powder will adhere and may be melted in the flame. In this way powder is to be constantly added. until a drop or bead is obtained which completely fills the loop. This bead must be perfectly colourless and transparent when it

is cold.

If the melted brad easily drops from the wire, either the loop or the bead is too large. A loop which is somewhat less than an eighth of an inch (3 mm.) across answers well; it may be made by rolling the tip of the wire round a piece of stout wire. If the bead is too large, remove a portion of it by giving the wire a slight jerk while the borax is in a fused condition.

If the bead thus formed shows any colour, it is ierked off while * is in a fused condition, and another bead is made from fresh borax. This process is repeated, if necessary, until a bead is obtained, which is colourless after it has been heated in the outer blowpipe-flame.

EXPERIMENT 27,--Moisten the clear cold borax-bend, and dip it into some finely-powdered manganese dioxide; a few minute particles only are to be thus attached to the bead. Then fuse this substance into the bead, by heating it for a short time in the tip of the outer blowpipe-flame.

Examine the colour of the bead at once by looking through it at a piece of white paper, or at a window-light or bright flame. It will be purple or amethyst-red while hot, and the colour will not alter when the bead has cooled.

Now heat the bead again for some time in the tip of the inner blowpipe flame. Its colour will vanish. The colour will, however, reappear when the bead is heated at the tip of the outer blowpipe flame. ,

These changes of colour in the borax bead are characteristic of the metal manganese and its compounds.

General Remarks on the Borax-bead .-- A beginner often obtains an opaque bead in the above experiment, because too much manganese oxide has been used. The bead should then be fused again, and a portion of it should be shaken off and replaced by fresh borax.

When the test with the borax-bead is finished, the bead is detached, while it is still fused, by giving the wire a sucklen jerk; or the bead may be removed, after it has cooled, by crushing it on a hard flat surface by a blow with the

pestle.

Two mounted platinum-wires should be kept, and their loops should always be immersed in dilute hydrochloric soid when the wires are not in use. They will then only require to be washed with water before they are employed for the above test.

59. The Microcosmic-bead.—Some substances give more decisive colorations to a fused bead of microcosmic salt (NaNH₄HPO₄4H₂O) in the platinum-wire loop, than to a bead of borax prepared as has been described in the preceding paragraph. The platinum-wire loop for this test must be made smaller than for borax, else the bead easily drops out of the loop.

60 Ignition on Charcoal in the Blowpipe-flame.—Wood charcoal is often employed as a support for substances which are to be strongly heated in the blowpipe-flame. It possesses the advantage of being cheap and infusible, and a bad conductor of heat.

Charcoal is, however, often used on account of the reducing or deoxidising power which it can exert at a high temperature. This renders it entirely unsuitable for oxidising processes such as that described in Exp. 25 (56).

Ignition on charcoal, therefore, serves chiefly to detect the presence of the heavier metals in their compounds. These are smelted out by the reducing action of the inner blowpipe-flame, aided by that of the red-hot charcoal. The assistance of fused reagents is, however, often necessary or at least advantageous.

The vapours of some metals, which have been thus smelted out, burn as they leave the charcoal, and the metallic oxides thus formed are deposited as "incrustations." The white ash, which is left by combustion of the charcoal, must not be mistaken for such an incrustation.

EXPERIMENT 28.—Choose a piece of wood charcoal which is free from large cracks, and which does not crackle or spit when it is heated in the blowpipe-flame; or, better still use a specially prepared hard charcoal block, which may be purchased and is more suitable and durable. Scoop out a *small* hollow near one end of it, with a knife or with a charcoal-borer.

Place in this cavity a small quantity of a mixture of powdered lead acetate and sodium carbonate. Then heat this mixture in the inner blowpipe-flame, holding the charcoal in such a way that the flame plays over its surface (Fig. 46, page 44).

Bright globules of lead will soon be visible in the cavity; and when the charcoal is removed from the flame, the surrounding surface will be found to be coloured with a yellow incrustation.

Take one of the little globules of metal from the charcoal with the point of a penknife, and strike it smartly with the pestle on the bottom of the inverted mortar. It will flatten out into a cake, showing that the metal is malleable and not brittle.

Detach another metallic globule, cleanse it from charcoal by, rolling it between the moistened finger-tips, transfix it on the

point of the penknife blade, and rub it on some paper. It will be found to mark the paper as a black-lead pencil does.

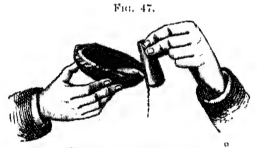
Lead compounds are characterised by giving a yellow incrustation, and midleable globules which are soft enough to-mark paper,



IGNITION ON CHARCOAL.

when they are subjected to this test on charcoal. This reaction therefore serves to detect the presence of lead in its compounds.

61. Washing and Levigation of the Residue left on the Charcoal.—The metallic globules, which have been obtained in the preceding experiment, are usually visible, even without the aid of a lens. They are also usually detachable from the charcoal, and do not require the following treatment.



WASHING BY LEVIGATION.

But fine metallic particles or scales can often only be separated and detected after the fused residue has cooled and been detached from the charcoal with the point of a penknife, and has then been washed by levigation. With this object, the detached residue is crushed in a mortar, or in a porcelain dish, with a little water. The water is then quickly decanted down the wet pestle, which is pressed against the edge of the vessel (Fig. 47). The light suspended particles of charcoal, together with soluble substances, are thus poured off; while the heavier and insoluble metallic particles are left in the vessel. By repeating this process of "levigation," the metal may be obtained free from soluble substances and from charcoal; it may then be further examined by a lens, or may be tested with a magnet or by other means.

It may be noted that the addition of a little potassium cyanide to the mixture, before it is heated on the charcoal, will usually much facilitate the process of smelting out the metal, and will assist in bringing it into the globular condition.

EXPERIMENT 29.—Fuse some sodium carbonate with a little copper sulphate on charcoal in the inner blowpipe-flame, and levigate the residue as is directed above. Little red scales of copper will be obtained.

EXPERIMENT 30.—Treat in the same way a mixture of ferrous sulphate and sodium carbonate: a grey powder, consisting of metallic iron, will remain. When this powder is touched under water with the point of a magnetised penknife, it will attach itself in the form of a tuft.

FLAME-COLORATIONS.

62. Characteristic Colours are imparted to a Non-luminous Flame by the vapours which many substances emit at a high temperature.

The blowpipe-flame may be employed to obtain these colorations.

A Bunsen-flame also answers the purpose well, but on account of the lower temperature of this flame, many colorations are not obtained with promptness or intensity.

When the Bunsen-flame is used for this purpose, it may be partly surrounded with a conical chimney, which steadies the flame considerably (Figs. 48, 49, 50). The supply of air, entering by the airholes of the burner, should be so regulated that the inner blue flame is sharply defined, and the platinum wire should be held either in the blue point or just below it.



BUNSEN-BURNER WITH CHIMNEY.

EXPERIMENT 31.—Heat a platinum wire loop, which is not more than an eighth of an inch (3 mm.) in diameter, in the inner blowpipe-

flame. If the wire is clean, it will become red hot without colouring the flame.

If the wire colours the flame, it must be boiled with hydrochloric acid in a test-tube and then rinsed with water. Or it may be wetted with strong hydrochloric acid, and then heated strongly in the blowpipe-flame until it no longer colours the flame. If the loop cannot be cleansed in this way, the end of the wire must be cut off, and a fresh loop must be formed.

Now dip the wire-loop into a little sodium chloride solution on a watch-glass, and remove it filled by a drop of the solution. Then hold the loop at the tip of the inner blowpipe-flame: the outer flame will be coloured bright yellow.

Repeat the experiment, placing the loop containing the solution in the outer part of the Bunsen-flame about one-third from the top. The same yellow tint will be seen above the loop.

This yellow coloration is given to the flame only by sodium and by its volatile compounds. On looking at it through the indigoprism (1369), no yellow coloration will be visible.

EXPERIMENT 32. Remove all the sodium chloride from the platinum wire loop by heating it strongly in the tip of the blowpipe-flame as long as it imparts any colour to the flame. Then dip the wire-loop into some solution of pure potassium nitrate. This will impart to the flame a pale lilac colour, which is characteristic of potassium and its compounds. When this coloration is viewed through the indigo-prism it will appear crimson.

EXPERIMENT 33.—Now dip the wire loop into a mixture of a few drops of the sodium chloride solution with a few drops of the potassium nitrate solution, which has been made on a watch-glass. When the loop is held in the flame the sodium coloration only will be seen. But if this coloration is then viewed through the indigoprism, the crimson potassium coloration will be seen distinctly. By means of the indigo-prism the potassium coloration is therefore discoverable, even when it is concealed by that of sodium.

THE SPECTROSCOPE.

63. The Spectra furnished by the Spectroscope afford the most certain means of detecting substances by means of their flame-colorations, more especially when the colorations due to several substances are mingled in the flame.

After the coloured light from the flame-coloration has passed through the slit of the spectroscope, it is separated into its constituent colours by traversing a glass prism.

The coloured bands or lines, which are seen as the result of this

separation, are termed the *spectrum* of the substance. By their colours and relative positions they serve to identify the substance with certainty: and this is true whether the flame-coloration is produced by one substance only or by several, since the constituent colours are seen without interference one with another.

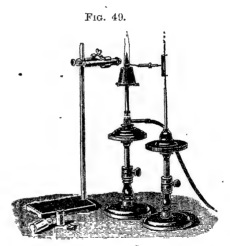
Two forms of spectroscope are in common use for chemical analysis.

The Direct-vision Spectroscope, which is shown ready for use in Fig. 66, is recommended by its compactness and its portability. In its smallest form it can be carried in the waistcoat pocket.

The Table Spectroscope, shown in Fig. 50 (page 48), is, however, to

be preferred for general use as a stationary piece of apparatus in the laboratory.

The light enters each of these instruments through a slit, the width of which can be adjusted to suit strong or feeble flame-colorations. The rays of light, after passing the slit, are rendered less divergent by traversing a lens, and then pass through one or more prisms. The rays are finally focussed by means of a sliding eye-piece, so as to give a sharp image of the slit.



THE DIRECT-VISION SPECTROSCOPE.

Note.—Spectroscopic observations should be made by looking at the flame against a black background: a piece of black velvet hung behind the flame serves well for this purpose. It is also advantageous to check in other ways the admission of extraneous light into the slit. Delicate observations should therefore be carried out in a more or less darkened room; but in ordinary analytical work the spectroscope may be used on the working bench in the laboratory.

64. The Direct-vision Spectroscope may be held by the fingers, or if it is not provided with its own stand, it may be

supported in a clamp in front of the flame (Fig. 49).

The slit is then slightly opened, and the eye-piece is adjusted until the vertical yellow line is sharply focussed. This line is always visible and is due to the sodium compounds present in the atmospheric dust which enters the flame.

If the spectroscope is held in the hand, it may be readily

focussed, before it is directed on the flame, by rendering sharp the dark Fraunhofer lines, which are seen on looking through a window at ordinary daylight.

More elaborate forms of the direct-vision spectroscope are available than that shown in Fig. 66. Most of these possess a scale for registering and mapping the position of the spectrum lines. If such an instrument is in common use, it is advisable to make out a spectrum chart, for purposes of reference, by the procedure which is described in paragraph 66.

Proceed to Experiment 34, attending to the Note (63).



THE TABLE SPECTROSCOPE.

55. The Table Spectroscope (Fig. 50) has usually three brass tubes. One of these carries the adjustable slit, another the eyepiece, and the third a transparent photographic scale.

The instrument is brought into adjustment by placing a Bunsen-burner with closed air-holes in front of the slit (Note, 63). On looking through the eyepiece at this luminous flame, a "continuous

spectrum" of unbroken colours will be seen. The upper and lower edges of this spectrum must be sharply focussed by moving the eye-piece, and the two tubes then arranged at such an angle as to give the longest complete spectrum possible. The tubes must either be permanently clamped in this position, or they must be so marked that they can at any time be easily brought into the same position with certainty,

Air is then admitted into the Bunsen-flame by opening the holes of the burner, and the yellow sodium line, which is always visible, is brought into a vertical position if necessary by rotating the slittube: the line is then sharply focussed.

The luminous flame is now lighted opposite the end of the scaletube, the scale is focussed, and the scale-tube is shifted until the yellow sodium-line stands at a particular graduation; the fiftieth graduation has been selected for the special chart in Fig. 51.

This yellow line will usually be seen in all the following experiments, since it is derived from the atmospheric dust, which enters

the flame. It may be produced, if necessary, with greater intensity by introducing a platinum wire, moistened with sodium chloride solution, into the flame.

EXPERIMENT 34.—When the spectroscope has been brought into adjustment, the student should proceed to introduce a mounted platinum wire (14) into the front of the Bunsen-flame, at a level just below the slit of the spectroscope. The platinum wire loop should be moistened successively with solution of sodium chloride, potassium chloride, barium chloride, strontium nitrate and calcium chloride.

The wire may be conveniently supported in the flame by slipping its glass handle upon a horizontal wire arm, the height of which is adjustable on the stand represented in Figs. 49, 50.

Coloured vertical lines will be seen when these flame-colorations are observed through the spectroscope. These lines will always be the same in colour and position for the same flame-coloration. They will vary, however, in these respects with the colorations of different substances. A particular "line-spectrum" is therefore characteristic of each of the vapours which give rise to the above flame-colorations.

66. The Spectrum-chart.—The small pocket-spectroscope is not usually furnished with a graduated scale for measuring the position of the spectrum-lines. In the absence of a scale the relative positions of the lines of any particular spectrum may, however, be jotted down or remembered with approximate accuracy. Since the yellow sodium line is never absent, it will furnish a useful landmark in the spectrum.

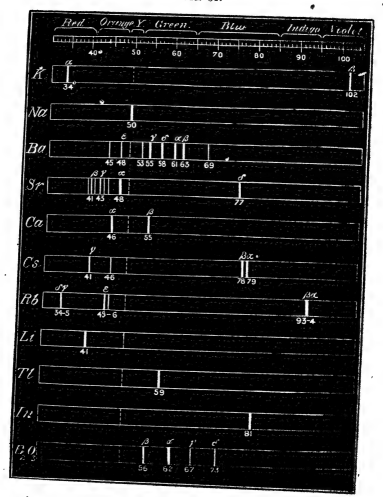
The scale of the larger spectroscopes, however, enables the position of the spectrum-lines to be noted down on a chart, and they are thus easily identified if the instrument is in adjustment. The instrument must always be brought to precisely the same adjustment as it possessed when the lines were mapped down in the spectrum-chart, if the chart is to be of any use.

The spectrum-lines will vary in position in different instruments, owing to variation in adjustment, and to difference in the material and angles of the prisms. Hence it is necessary to make a separate chart for each spectroscope.

The accompanying spectrum-chart (Fig. 51, p. 50) indicates a convenient method of mapping the lines, but the position of the lines on the scale only applies to the spectroscope from which this chart was made.

In the chart the relative intensity of the spectrum-lines may be inferred from their varying thickness, while the relative intensity of the lines in each spectrum is indicated by the Greek letters applied in the order of decreasing intensity.

Fig. 51.



THE SPECTRUM-CHART.

The colour of each line is shown by the name of the colour, which is placed at the top of the chart over the region in which the line occurs. The position of each line is indicated by placing t it the number of the scale-graduation at which i

reasons which have been already stated, and it therefore serves as a landmark.

A chart of this kind must be at hand for reference when the spectroscope is used for analytical work. The spectrum-lines may be marked in with pen or pencil on a white ground, but they are more striking if coloured. The chart may conveniently be suspended near the instrument.

The spectrum of an unknown substance may also be identified, even when the instrument has not been carefully adjusted, and in the absence of the chart. For this purpose the position of the lines is noted on the scale and the flame-coloration of a known substance, which is believed to correspond to the spectrum which is under observation, is then thrown into the spectroscope. If the spectrum-lines of the unknown substance are identical in colour and in position with those given by the known coloration, the two spectra must be due to the same substance, and the unknown substance is therefore identified.

EXPERIMENT 35.—The student should carefully observe and map the spectra of several of the metals, which are shown in the spectrum-chart (Fig. 51). He should then proceed to examine some of the mixed spectra, which are obtained from solutions containing the salts of two or more of these metals. The ease with which many metals can be found by means of the spectroscope, even in the presence of one another, will thus become evident.

The Delicacy of this Method of detecting certain metals is extreme. Thus it is only in air which has remained undisturbed for a long time and is almost absolutely free from dust particles, that the sodium spectrum is not obtained from a Bunsen-flame. By using extremely dilute solutions of metallic salts, the delicacy of the method has been proved in the case of other metals. Bunsen states that one three-thousand-millionth of a grain of sodium can be detected by the spectroscope.

In the more Powerful Spectroscopes several prisms are used, and the lines of the spectra become more widely separated. Their relative positions are noted with great precision by moving the observing-tube until each line in succession is brought into the middle point of the cross-wires in the eyepiece. The position of the eye-piece is then read off on the graduated edge of the circular table on which the prisms stand, and registers the position of each line. Instruments of great power and precision are, however, not requisite for ordinary analytical work, and the simple form of table-spectroscope which has been already described will be found to be sufficient for the purpose.

- 67. Use of the Spectroscope in Chemical Analysis.—The spectroscope is of special use in chemical analysis for the following purposes:—
 - 1. For readily detecting metals when their flame-colorations

are concealed by that of sodium; and generally for detecting metals, when their colorations are mixed in the flame.

- 2. For distinguishing similar colorations from one another. Thus the red coloration of strontium is easily distinguished from that of calcium and of lithium, by the strongest red line in each of the spectra being at a different distance from the sodium line; these very similar flame-colorations are further distinguished by the appearance of a characteristic blue line in the spectrum of strontium, and by a characteristic green line in the spectrum of calcium.
- 3. For detecting minute traces of many metals, which could scarcely be discovered by other means. The metals lithium, indium, thallium, rubidium, and essium were discovered by their spectra; and as these metals usually occur in minute proportion only, they are most conveniently detected by the spectroscope.

It is by no means a matter of indifference which compound of a metal is selected for obtaining the flame-coloration or spectrum of the metal. The most volatile salts are generally the chloride, the nitrate, and the chlorate; these accordingly give the most vivid but the least durable spectra. The coloration is increasingly feeble with the sulphate, the carbonate, and the phosphate; and the silicate does not usually give any coloration at all.

The coloration becomes more intense when the finely powdered substance is moistened with strong hydrochloric acid. But in the case of a silicate it is usually necessary to moisten the powder with hydrofluoric acid, or to mix it with pure ammonium chloride and calcium carbonate, before it is introduced into the flame.

Some very volatile substances give only a momentary flash of colour in the flame; while others, which are far less volatile, colour the flame only at the highest temperature obtainable. Hence careful observation of the flame-coloration through the spectroscope is necessary, from the moment that the substance is introduced into the flame, until the wire-loop has been heated to whiteness for some time.

The width of the slit of the spectroscope should be regulated according to the intensity of the lines to be looked for. A bright line can be observed with a narrow slit, and its position is thus noted with great precision. The slit must, however, be opened much more widely to see a faint line, such as $K\beta$ (Fig. 68). It is undesirable that the lines should be of great breadth, since breadth prevents the position of lines on the scale from being accurately noted, and may even cause the edges of neighbouring lines to overlap.

PART II

ANALYTICAL REACTIONS

INTRODUCTORY REMARKS.

80. Object and Method of Chemical Analysis.—The study of chemistry has revealed the fact that every substance consists either of one kind of matter only, or of two or more different kinds.

Those substances which contain only one kind of matter are called "Chemical Elements." Their number at present amounts to about eighty. A list, which comprises most of them, is given at the end of the book (1420).

These elements do not usually occur singly. Two or more of them are commonly found together either in a state of mixture, or united in a more intimate manner in a "chemical compound."

It is the object of analytical chemistry to discover what element, elements, or sometimes what groups of elements or "radicles," any substance of unknown composition contains. But an analysis may also be undertaken in order to ascertain whether any particular element or group of elements is present in a given substance.

An analyst should be acquainted with the properties of each element and of the compounds which it forms with other elements. Efficiency in chemical analysis therefore presupposes a knowledge of what is usually termed Descriptive or Theoretical Chemistry.

For ordinary analytical purposes, however, certain marked properties or changes of each element, or group of elements, are selected; and by these properties, called "tests" or "reactions," its presence is detected.

Substances, which are added for analytical purposes, are called "reagents." Some reagents serve for detecting one element or compound only, and are therefore called "special reagents"; others separate or detect a group of elements or compounds, and are called "group reagents" or general reagents.

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Two classes of reactions are employed in qualitative analysis:—
(1) "Dry Reactions" which are applied in the solid state. Some dry tests have been already described. As a rule these tests are more limited in their application and less delicate than the wet reactions, but since they do not require elaborate or fragile apparatus, they can be applied when a laboratory is inaccessible. Dry tests are therefore commonly used when one is prospecting for minerals and ores in comparatively remote places.

(2) "Wet Reactions."—The reactions usually employed in chemical analysis are, however, applied in solution, since wet tests are usually more delicate, and more generally applicable.

A short account of theoretical considerations involved in the use

of wet reactions is given below.

81. Analytical Components of Dissolved Substances.—Salts, acids and bases, when in solution, do not behave as complete molecules, but as if they have been resolved into simpler components termed "ions."

Thus when hydrochloric acid (HCl) is dissolved in water, it is largely resolved into H-ions and Cl-ions. A similar "ionisation" occurs in the case of other soluble acids; and it is generally true that the solution of a compound, which produces H-ions when it is dissolved in water, gives an acid reaction to litmus.

The solution of an alkali, such as potassium hydroxide (KOH), contains the ions K and OH, the latter being the ion which gives the alkaline reaction to litmus.

The extent to which the molecules of an acid or base undergo ionisation when they are dissolved varies considerably with different substances and solvents. Strong acids, such as hydrochloric sulphuric and nitric acids, when they are dissolved in water undergo a high degree of ionisation; whereas weak acids, such as carbonic nitrous and boric acids, are only slightly ionised by water. Similarly the powerful bases, potassium and sodium hydroxides, exist in aqueous solution largely in an ionised condition; whereas the weaker ammonium hydroxide is ionised in water to a much smaller extent.

The solutions of many salts, such as potassium chloride (KCl), are neutral to test papers, since their solutions contain neither H (acid) nor OH (basic) ions. These salts may usually be produced by the action of the powerful acids upon the strong bases.

A large number of salt solutions, however, change the colour of test papers; the majority of them showing an acid reaction, and a lesser number yielding an alkaline reaction. In these solutions the acidity and alkalinity are due to H and OH ions, which have been produced by a further reaction termed "hydrolysis." The solution

of a compound formed by the reaction of a strong acid upon a weak base usually gives an acid reaction, whereas the solution of the compound similarly produced from a weak acid and a strong base usually gives an alkaline reaction.

For a fuller explanation of the theory of ionisation the student is referred to works on chemical theory.

Assuming that the resolution into component ions occurs when salts are dissolved in water, the general applicability of the test for an element or radicle to a series of salts becomes intelligible. Thus a solution of any salt of calcium will give a white precipitate of calcium carbonate with ammonium carbonate, since the metal s separated by solution and acts independently of the other portion of the salt. Similarly any soluble metallic chloride will give white precipitate with silver nitrate solution, because the ionised acid or chloride portion of the salt acts independently of the metal.

The systematic name of a salt or acid is given in accordance rith the above considerations and consists of the name of the netal or hydrogen which the compound contains; associated with he name of the element or radicle with which the metal or hydrogen combined.

Thus in the case of sodium sulphate, the name of the metal sodium" is associated with that of the "sulphate" acid-radicle, ad when this salt is dissolved in water the ions present in solution to Na and SO₄. The latter portion of the name of such a chemal compound usually denotes an acid-radicle, but occasionally enotes hydroxyl or oxygen or some other element.

Similarly acids are frequently called hydrogen salts, because a property of the second
From the above examples it will be seen that some of the aciddicles are elements, whilst most of them are "compound radicles."
The above division of the name of a chemical compound, generally
responds to the portions of the compound substance which are
parately detected during its analysis. For analytical purposes
stances are accordingly classed as Metals and Acid-radicles.
Dertain elementary and compound substances cannot, however,
included in this generally useful classification; and certain nontallic or metalloidal substances, as well as certain acid-radicles,
for analytical purposes conveniently classed with the metals.

82. Analytical Groups.—The metals are separated, by their behaviour with certain general reagents, into five principal Groups, and two of these Groups are further subdivided into two Subgroups.

This classification is tabulated on the opposite page.

Each Group receives a distinctive name, which is derived either from its place in the system, or from some conspicuous or important member of the Group, or from the general reagent which is used to precipitate the Group. Thus we speak indifferently of the Fourth Group, of the Barium Group, or of the Ammonium Carbonate Group.

It is perhaps well to avoid referring to a Group by its number, since different analysts number the groups differently, and the *number may not therefore denote the Group with certainty.

As will be noted later on (462), the acid-radicles cannot be as satisfactorily grouped for analytical purposes as the metals, since they do not usually undergo complete precipitation in groups the members of which admit of being subsequently separated from one another.

TRYING AND RECORDING THE ANALYTICAL REACTIONS.

83. In order to become fully acquainted with the behaviour of different substances with reagents, and thus to be able to detect these substances by their reactions, each of the reactions should be tried with the substance itself or with one of its compounds. The appearances which are presented are carefully observed and remembered, and the substance may then be easily recognised in a body of unknown composition by means of these reactions or tests:

The addition of a reagent to a substance causes some important and visible change to occur. This may be simply a change of colour, but it very usually consists in the formation of a precipitate or insoluble substance in the solution. The colour, the appearance, or the behaviour of this precipitate with other reagents, is characteristic of a particular element or compound-radicle.

As soon as a reaction has been satisfactorily performed, it should be briefly described in the notebook in a neat and concise manner. The nature of the chemical change which has occurred should then be explained by means of a chemical equation. Generally the use of chemical symbols and formulæ instead of chemical names may be adopted for purposes of brevity, and the use of the following abbreviated formulæ and words will also be advantageous.

ANALYTICAL CLASSIFICATION OF THE METALS.

By adding each of these Group-reagents in the above order to the filtrate or clear solution from the preceding Group-precipitate, the metals in solution may therefore be separated from one another into five groups.

Lead will be found both in Groups I, and II., since its chloride is somewhat soluble.

84. Abbreviations employed for Entry in the Notebook.—For convenience in hasty writing, it is not uncommon to employ abbreviated formulæ for the compound radicles. This is more especially the case with those radicles which are constantly met with, and with those which have a complex composition. Only such abbreviations are employed in the text as save the use of brackets or of numbers. Thus Am_2 is short for $(NH_4)_2$, and $(!y_2)_2$ for $(!y_2)_2$ or $(!x_2)_2$.

The following abbreviations may be used:-

Am for (NH₄), the metallic-radicle ammonium.

 \overline{T} ,, $(C_4H_4O_6)$, the acid-radicle of tartrates.

 $\overline{4}$,, $(C_2H_3O_2)$, the acid-radicle of acetates.

 \overline{Ci} ,, $(C_6H_5O_7)$, the acid-radicle of citrates.

Cy ,, (CN), the acid-radicle in cyanides.

The following contracted words may also be used:-

	expt.	for	experiment.	insol.	for	insoluble.
	pp.	,,	precipitate.	sol^n .	**	solution.
	pp^n .	"	precipitation.	cryst.	,,	crystalline.
1	pp^d .	,,	precipitated.	col^n .	,,	coloration.
1	sol.	,,	soluble.			

85. Example of Entry of Reactions in the Notebook.—The following example, selected from paragraphs 108-113, will serve as a specimen of the brief entry of reactions in the Note-book.

SILVER.—Used AgNO₃-solution.

Test.	Observation.
109. K ₂ CrO ₄ 112. AmOH 113. Mixed solid with Na ₂ CO ₃ and heated on charcoal if the inner blowpipe-flame.	White pp., became curdy when heated; sol. in AmOH and in KCN, and repp ^d on adding HNO ₃ in excess; insol. in hot HNO ₃ ; darkened by exposure to light: HCl + AgNO ₃ = AgCl + HNO ₃ . Deep crimson pp.: K ₂ CrO ₄ + 2AgNO ₃ = Ag ₂ CrO ₄ + 2KNO ₃ . Light brown pp. on adding a little dilute AmOH, easily sol. in excess:— AmOH + AgNO ₃ = AgOH + AmNO ₃ . White metallic bead obtained, which was malleable but did not mark paper: no increastation formed.

GENERAL RULES FOR WORKING.

- 86. The student should attend carefully to the following precautions. They will materially aid the progress and accuracy of his work, and if they are observed from the commencement they will not be felt irksome.
- 1. Before beginning work, look through the reagent-bottles belonging to the working bench (1409). Replace any which are absent, and see that the bottles are arranged in a convenient order. Then proceed to fill any which are empty, and to relabel any whose labels are imperfect or loose.
- 2. Arrange the apparatus, which is required for use, on the back part of the bench, so as to leave the front of the bench free to work upon. This will lessen the risk of upsetting anything. (See Fig. 66, p. 331.)
- 3. All dirty glass and porcelain, which is not in use, should then be carefully cleansed. This ought, however, to be unnecessary, since apparatus should never be put away dirty.
- 4. When a reagent-bottle is to be used, take it in the right hand; remove the stopper by grasping it between two of the fingers, or between the fourth finger and palm of the left hand; hold the stopper in this way while the bottle is being used; then replace it immediately, and return the bottle to its place. Do not lay down the stopper on the bench.
- 5. Pour a liquid reagent out of the side of the bottle opposite to that which bears the label; and prevent the last drop of the liquid from running down the outside, by touching the lip of the bottle either with the stopper, or against the clean edge of the vessel.
- 6. Solid reagents should be taken out of the bottle on a clean knife-blade or spatula, or with a slip of clean cardboard or paper.
- 7. Each bottle must be restoppered and replaced on the shelf, in its proper place and with the label outwards, immediately after use; it must never be left standing on the working-bench.
- 8. If a solution or precipitate has to be put by for some time before it undergoes further examination, it should always be labelled. A piece of gummed paper may be attached to it, bearing either an inscription, or a reference number which is explained in the note-book and which is sufficient to recall to mind what the solution or precipitate consists of. It is well never to trust to memory in this matter.

It will be found to be unadvisable as a rule to leave a precipitate

thus, since it is apt to dry or to undergo oxidation.

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- 9. It should be remembered that if metal crucible-tongs are employed for holding hot evaporating basins containing acids, some of the metal is apt to be dissolved and thus introduced into the solution. A hot dish may be moved either by holding it with a cloth, or by lifting the stand upon which the dish is placed.
- 10. Liquids only should be poured down the sinks. All solid refuse, such as soiled filters, broken glass, and pieces of charcoal, should be thrown into a receptacle which is placed in an accessible position.
- 11. When a substance is being examined in order to find out its composition, the student should fully write down in the note-book each test or process, with the result obtained, as soon as it is completed. The analysis is thus written out gradually as it progresses. On no account should the entry of the results be left until the completion of the analysis.

This rule cannot be too strongly enforced, as neglect of it always causes uncertainty and confusion. The most convenient form of entry is that of the Analytical Tables.

12. Cleanliness and neatness in analytical work cannot be too carefully attended to. More confusion and error may be caused by using dirty test-tubes, beakers, dishes, and funnels, than would be thought possible by the beginner. Work should also be rendered as systematic as possible, in order to economise time and thought.

87. Directions for Trying the Analytical Reactions:

1. A solution of the substance, the reactions of which are to be tried, must always be employed unless it is specially stated that the solid is required. The solid substance is usually only required for blowpipe reactions.

A solution may be kept in readiness, or it can be easily made by warming the solid with water or with dilute acid (1411-1413).

- 2. Pour into a test-tube or small beaker about twenty cubic centimetres (20 c.c.) of the solution of the substance: this constitutes the "original solution." If the solid substance is also required, take some of it upon a watch-glass and powder it finely.
- 3. Pour about 2 c.c. of the original solution into a perfectly clean test-tube and add the first-mentioned reagent. A few drops only of the reagent should be added at first, more can be gradually added afterwards if necessary.

The habit of working with small quantities both of solutions and of reagents must be acquired from the first.

In trying each of the reactions a separate portion of the original solution is to be taken.

4. Before trying a reaction, the student should carefully read through the whole of the description of the results to be obtained. The experiment is then to be performed, literally following out the directions given.

It is a necessary condition of after success that each result should be obtained precisely as is stated in the text. The student must never on any account pass on without being satisfied that the statements of the book are true, and that he could at any time repeat the test successfully.

5. The water used for the tests and reactions must always be distilled water.

The acids are to be used in a dilute condition, unless it is stated that the strong acid is to be employed.

In all cases KOH and NaOH may be used indifferently the one for the other.

SECTION III

REACTIONS FOR THE METALS

INTRODUCTORY REMARKS.

99. Paragraphs which are printed in small type may be passed over or simply read through if time is limited; they should, however, receive careful attention from the general analyst.

Since chemical symbols and formulæ are freely used in the text, both the name and the symbol or formula for each reagent are given in the following reactions when the reagent is first mentioned; subsequently the chemical formula only is used. Any difficulty in connecting symbols and formulæ with chemical names should also be prevented by referring to the tables giving the names and corresponding symbols of the chemical elements (1420), the formulæ of minerals containing them (1419), and of the substances used for analytical purposes (1409-1418).

The list of minerals containing the more commonly occurring metals (1419) furnishes information as to the compounds in which these metals occur in nature, and also indicates by formulæ the chemical composition of these natural compounds.

The chemicals and reagents required for the work of Parts II and III of the book will be found tabulated in Part IV (1407-1414), where also their preparation is described (1383-1400).

Occasionally non-metallic or metalloidal elements are associated with the Metal Groups for the purpose of analytical classification.

The reactions of the Rarer Metals belonging to each Group are entered in small type at the end of the Group.

GROUP I.—SILVER GROUP.

100. This Group includes Pb, Ag, Hg, together with the rares elements Tl and W.

The solutions of Pb, Ag, Hg, Tl differ from those of all other metals, by being pracipitated as chlorides by hydrochloric acid (HCl) or by a dissolved chloride. Since, however, lead chloride (PbCl) and thallium chloride (TICl) are somewhat soluble in water, these metals are not entirely precipitated in this Group as chlorides, and they will also be found in Group II.A. as sulphides in the course of analysis. W is precipitated in the course of analysis.

For the Rarez Elements in this Group refer to paragrants in 131.

Lead (Pb).

Density 11.37, melting point 327.4° C.

• IOI. Lead is a bluish-white metal, which slowly tarnishes in the air, and is so soft that it may be indented with the finger nail, and will readily yield a streak on paper. It is insoluble in dilute sulphuric and hydrochloric acids, but dissolves in nitric acid with the evolution of nitric oxide.

Lead forms the following well-known oxides, lead monoxide (PbO) yellow, red lead (Pb₃O₄), and lead peroxide (PbO₂) brown.

The lead salts correspond to the monoxide. The lead-ion is divalent and colourless.

For lead-minerals refer to par. 1419.

REACTIONS OF LEAD.

Use Lead acetate, PbA2.3H2O, dissolved in dilute acetic acid.

- ro2. Hydrochloric acid (HCl): white precipitate (PbCl₂) which forms only in cold and strong solutions. Boil the precipitate with sufficient water: it will dissolve, but the PbCl₂ will separate again in slender crystals, when the hot solution is slowly cooled.
- 103. Hydrogen sulphide (H₂S): black precipitate (PbS). This precipitate is often red if much hydrochloric acid is present in the solution; but the red colour changes to black when water is added and H₂S is passed, or when H₂S-water is added.

Filter, and show with separate portions of the precipitate that PbS is insoluble in potassium hydroxide (KOH) and in ammonium sulphide (Am $_2$ S), but is soluble in boiling dilute nitric acid (HNO $_3$); also that it is converted by boiling strong HNO $_3$ into white insoluble PbSO $_4$.

104. Sulphuric acid (H₂SO₄): white precipitate (PbSO₄). This precipitate is much less soluble in dilute H₂SO₄ and in alcohol than in water; hence the dilute H₂SO₄ should be added in excess to expretty strong solution of the lead salt.

Shape up the liquid, and pour it off into two test-tubes; let the liquid stand in each of these tubes and decant it from the precipitate (42). Show that the precipitate may be dissolved by boiling it either with ammonium acetate (AmA) solution, or with acetic acid (HA) after adding ammonium hydroxide (AmOH) in excess; show that it dissolves in boiling strong HCl.

Potassium chromate (K2CrO4): yellow precipitate (Phron Pour off the liquid with the precipitate into two tubes,

and show that the precipitate is soluble in KOH, but insoluble in HA.

106. Blowpipe-test on Charcoal.—Mix some powdered lead acetate with about twice as much powdered sodium carbonate (Na₂CO₃) in a cavity on a piece of charcoal, and heat the mixture in the inner blowpipe-flame (Exp. 28, 60).

A yellow incrustation will form upon the surface of the charcoal around the mixture, and small lustrous globules of Pb will be seen within the cavity. The incrustation is readily removed by heating it in the inner blowpipe-flame, and it colours the flame meanwhile pale blue, unless this colour is masked by the sodium yellow.

Detach one of the metallic globules with the point of a penknife, place it on the flat bottom of an inverted mortar, and strike the globule a blow with the pestle. It will flatten out without cracking or breaking into powder, showing that Pb is $m \cdot ll^{ev}bl^{\nu}$ and not brittle.

If one of the cleansed globules is fixed upon the point of a penknife, and is then rubbed with gentle friction on paper, it will mark the paper as a blacklead pencil does.

Silver (Ag).

Density 10.47, melting point 960° C.

107. Silver is a white lustrous metal; it is highly malleable and ductile, and is not affected by air unless hydrogen sulphide is present, when it is blackened by the formation of silver sulphide.

Silver is insoluble in hydrochloric acid, but dissolves readily in nitric acid and in hot concentrated sulphuric acid. One series of salts of silver is known, which correspond to the monoxide (Ag₂O). The silver-ion is monovalent and colourless.

For silver-minerals refer to par. 1419.

REACTIONS OF SILVER.

Use Silver nitrate, AgNO₃.

108. HCl: white precipitate (AgCl), which when it is shaken or heated.

Pour off the liquid with the precipitate into four test-tubes. Filter one portion, then wash the precipitate and let it stand for some time exposed on the filter to sunlight or to common daylight; it will become blackish-purple.

Show with the other portions that AgCl dissolves readily when is heated with AmOH or with potassium cyanide (KCN)-solution,

and is precipitated again from these solutions when HNO₃ is added in excess. Show also that AgCl is insoluble in HNO₃, even when the acid is boiled.

- 109. K2CrO4: deep crimson precipitate of silver chromate, Ag2CrO4.
- 110. H₂S or Am₂S: black precipitate (Ag₂S), insoluble in Am₂S and in KOH, soluble in boiling dilute HNO₃.
 - III. KOH: brown precipitate (AgOH), insoluble in excess.
- II2. AmOH: light brown precipitate (AgOH), produced only when very dilute AmOH is added drop by drop, since AgOH is very easily soluble in excess of AmOH.
- II3. Blowpipe-test on Charcoal.—If any solid substance containing Ag is mixed with powdered Na₂CO₃ and the mixture is then heated on charcoal in the inner blowpipe-flame, white malleable globules or scales of Ag will be obtained, but no incrustation will be formed on the charcoal. The globules do not mark paper (106).

Mercury (Hg).

Density 13.6, melting point -39° C, boiling point 357° C.

114. Mercury is a silver-white liquid metal which remains unaltered in the air at ordinary temperatures.

The metal is insoluble in hydrochloric acid, but dissolves in hot concentrated sulphuric acid. Dilute nitric acid acts upon it slowly at ordinary temperatures, forming mercurous nitrate: the concentrated acid acts rapidly, forming mercuric nitrate, and coloured oxides of nitrogen are evolved.

Mercury forms two series of compounds: "mercurous compounds," corresponding to mercurous oxide Hg₂O, with a monovalent colourless metallic ion; and "mercuric compounds," corresponding to mercuric oxide, HgO, with a divalent colourless metallic ion.

The metal present in these two classes of compounds may be distinguished as "mercurosum" (Hg') and "mercuricum" (Hg') respectively.

Mercurosum and mercuricum have to be placed in different analytical groups, and their common reactions are not repeated in each group. The reactions which are given below are characteristic of mercurous salts and are distinctive of the two classes of salts, while those given under mercuric salts (134-136, 140 and 141) for common to both classes of salts.

Mercury occurs in pature as Cinnabar, HgS.

REACTIONS OF MERCUROSUM (Hg).

Use Mercurous nitrate, Hg₂(NO₃)₂.H₂O, dissolved in dilute HNO₃.

115. HCl: a white precipitate (Hg₂Cl₂), insoluble in cold dilute acids, but soluble when it is warmed with HNO₃ or with a mixture of HCl and HNO₃ which converts it into soluble mercuric chloride (HgCl₂).

This precipitate becomes black when AmOH is poured upon it, but does not dissolve: Hg₂Cl₂ + 2NH₄OH = NH₂.Hg₂Cl +

 $NH_4Cl + 2H_2O.$ $^{\circ}$

A solution of mercuric salt is not precipitated by HCI or by solution of a chloride.

- 116. AmOH, or KOH: black precipitate, insoluble in excess. In a solution of mercuric salt AmOH yields a white precipitate (137), and KOH gives a yellow precipitate (138).
- 117. Potassium iodide (KI): yellowish-green precipitate of $\mathrm{Hg}_2\mathrm{I}_2$, insoluble in excess.

In a solution of mercuric salt, KI produces a pink precipitate, which turns to scarlet, and is soluble in excess of either KI-solution or of mercuric solution.

DETECTION OF METALS IN GROUP I.

- · After the student has carefully worked through the reactions for metals in Group I., he should proceed to apply the knowledge which he has acquired to the detection of the metals belonging to this group, occurring either singly or together as is directed below.
- 118. A solution, which is to be examined for Pb, Ag, and Hg, is precipitated by adding HCl in excess (see *Note*) and the precipitate is examined by Table I. (1250).

Examination of a white solid for Metals of Group I.

The substance was dissolved in water, and dilute HCl was added in excess; the liquid was then filtered and the precipitate was washed several times with hot water, the washings being kept separate:—

Washings: a few white crystalline needles deposited	Residue: rinsed with AmOH into a test-tube and boiled, then filtered:—	
on cooling. Confirmed by adding \$\xi_2\text{CrO}_4\$, a copious yellow precipitate showed:—	Filtrate: added HNO ₃ in excess, no white precipitate formed:—	Residue on the filter was black:—
Presence of Pb.	Absence of Ag.	Presence of Hg.

Found Pb and Hg.

REACTIONS FOR RARER ELEMENTS IN GROUP I.

120. The rarer elements included in this Group are Tl and W.

The addition of HCl in excess partially precipitates Tl as chloride, and

completely precipitates tungstic acid from a tungstate.

Since TICl is only partially precipitated in Group I., owing to its being somewhat soluble in water, Tl is also found in the ordinary course of analysis in Group III.B., where it is completely precipitated as sulphide by Am₂S.

THALLIUM (Tl).—Use Tl₂SO₄-solution.

121. TI occurs in small quantity in many natural sulphides, and occasionally in the ashes of plants and in mineral waters. It is not a constituent of any characteristic mineral, and the principal source from which it is prepared is the dust which settles in the flues of the burners of sulphuric acid chambers, when pyrites is used in the manufacture.

Tl yields both thallic and thallious salts, but the former are very unstable,

changing, even when their solutions are heated, into thallious salts.

122. HCl: a white precipitate (TICl), which rapidly settles, does not blacken in the light, and is soluble in aqua regia. It is soluble in a large quantity of water, and therefore does not form in dilute solutions.

123. KI: a yellow precipitate (TII), almost insoluble in water, but more soluble in KI-solution. If ferric salt is present in the solution, it must be reduced by H₂SO₃ before KI is added. This is the most delicate test for TI.

• 124. H₂PtCl₆ (Note, 420): yellow precipitate (Tl₂PtCl₆), almost insoluble in water.

125. Am₂S: black precipitate, which is easily coagulated by heat; is insoluble in AmOH, in alkaline sulphides, and in KCN-solution; is readily oxidised by the air to Tl₂SO₄, and is easily soluble in mineral acids.

TI is completely precipitated by H₂S from a solution in which HA is the only free acid present, but the presence of a free mineral acid prevents the precipitation.

126. Flame-coloration.—Thallium compounds impart to the Bunson flame an intense green colour, which rapidly disappears.

The spectrum (Fig. 51, page 50) consists of one bright emerald-green line, and is very characteristic. Thallium may be easily detected by the spectroscope in solution, or in any of the precipitates which are mentioned above.

Tungstate (WO₄).—Use Na₂WO₄-solution.

- 127. W usually occurs native in the form of a tungstate. An insoluble tungstate yields a soluble alkaline tungstate when it is fused with an alkaline carbonate. From the solution of an alkaline tungstate HCl precipitates the tungstic acid completely.
- 128. HCl, HNO₃, or H₂SO₄: white precipitate of hydrated H₂W()₄ which becomes yellow and anhydrous when the liquid is boiled; it is insoluble in excess of acid, but soluble in AmOH.

If a piece of Zn is dropped into the acid liquid containing the precipitate, it yields a deep blue colour by reduction of the tungstic acid to W_2O_5 .

- 129. SnCl₂ in neutral solution, made by dissolving SnCl₂-crystals in water and filtering, gives a *yellow* precipitate, which becomes *blue* when H(! is added and the liquid is heated. This is a most delicate test for a tungstate.
- 130. Am_2S yields no precipitate in the solution of an alkaline tungstate; but if Am_2S is added, and the liquid is then made acid with HCl, brown WS_3 is precipitated.
- 131. Microcosmic-bead: outer flame, colourless or yellow; inner flame, blue; if a little FeSO₄ is heated with the fused bead, the colour changes to blood-red. These colours are best seen when the bead is perfectly cold.

For the Detection of the Rarer Elements of this Group, refer to the first column in the Table (1323).

GROUP II.—COPPER AND ARSENIC GROUP.

132. This Group includes the metals Hg, Pb, Bi, Cu, Cd, As, Sb, Sn, together with the rarer elements Pd, Os, Rh, Ru, Au, Pt, Mo, Se, Te, Ir.

The reactions of the rarer elements are given in paragraphs 210-261.

Group II. is subdivided into Groups II.A. and II.B.

GROUP II.A.—COPPER GROUP.

133. This Group includes Hg, Pb, Bi, Cu, Cd, (Pd, Os.)

The compounds of these metals differ from those of the metals in Groups III.A., III.B., IV., and V. by being precipitated as sulphides by H2S from their acid solutions. With the exception of Pb, which is partially precipitated from strong solutions as chloride, these metals are not precipitated from their solutions by HCl or by dissolved chlorides.

The precipitated sulphides of the metals of Group II.a. differ from those

of Group II.B. by being insoluble both in Am2S and in KOH.

H2S does not precipitate the members of this Group from strongly acid solution readily or completely; these solutions must therefore be suitably diluted with water in order to ensure their precipitation.

Mercury (Hg).

A statement of the general properties of this metal and its com-

pounds has already been given in paragraph 114.

It is also explained in that paragraph that mercury forms two classes of salts, the metal being termed mercuricum in the mercuric compounds which are classed in Group II.A., and mercurosum in mercurous compounds which are classed in Group I.

• The reactions given in paragraphs 115-117 and 137-139 are distinctive tests for mercury in these two classes of salts.

REACTIONS FOR MERCURICUM (Hg").

Use Mercuric chloride (HgCl₂).

134. H₂S: black precipitate (HgS). If the H₂S-solution is added slowly, the precipitate is first white, then brown or orange,

and ultimately black. These changes of colour, which are due to the formation of an intermediate compound (HgCl₂.2HgS), are characteristic of mercuric salts.

Filter off the precipitate of HgS, and wash it free from dissolved

chlorides; then place portions of it in two test-tubes.

Boil one of these portions with strong HNO₃, the precipitate will not be dissolved; now add HCl to this portion and warm again, the precipitate will dissolve.

Boil the second portion with Am2S, the precipitate will not

dissolve.

Hence HgS is insoluble in hot strong HNO₃, and in Am₂S. It is, however, easily soluble in a mixture of HNO₃ and HCl, which is known as aqua regia.

135. Stannous chloride, SnCl₂; white precipitate (Hg₂Cl₂): if an excess of SnCl₂ is added, the white precipitate becomes grey; examination with a lens shows that the grey precipitate consists of small globules of Hg:—

$$2 \text{HgCl}_2 + \text{SnCl}_2 = \text{Hg}_2 \text{Cl}_2 + \text{SnCl}_4$$

 $\text{Hg}_2 \text{Cl}_2 + \text{SnCl}_2 = 2 \text{Hg} + \text{SpCl}_4$.

Boil the liquid, and decant it from the grey precipitate. Then heat the precipitate with strong HCl; the Hg-particles will usually unite to form larger globules, which are visible through a lens or even to the naked eye.

136. Copper, Cu: cleanse a small strip of copper sheet, or a copper coin, by rubbing it with emery-paper, or by dipping it into strong HNO₃. Immerse it in some HgCl₂-solution, which has been acidified by the addition of a few drops of HCl. The Cu will become coated with a grey film of Hg:—

$$\sqrt{\text{HgCl}_2 + \text{Cu} = \text{Hg} + \text{CuCl}_2}$$

Allow the Cu to remain immersed in the liquid for several minutes, then remove it, dry its surface, and rub it with a cloth or with a piece of wash-leather. The Cu will appear more or less whitened like silver, since the mercury has formed an "analgam" with it.

The Hg may be readily driven off from the copper by heating it strongly. If the copper strip is heated in a test-tube, or better in a small glass tube sealed at one end (13), a sublimate of globules of Hg will form upon the cool sides of the tube.

137. AmOH: white precipitate of NH HgCl, insoluble in excess.

138. KOH: yellow precipitate of mercinic oxide (HgO).

139. KI: red precipitate of mercuric iodide (HgI₂), readily soluble in excess of either KI-solution or HgCl₂-solution.

140. Mercury mirror.—Dry some sodium carbonate perfectly by heating it strongly on platinum foil or in a porcelain dish. Also dry the interior of a small test-tube by heating it in the flame, and then sucking out the moist air by a glass tube, which has been pushed down to the bottom.

Mix a little finely-powdered HgCl₂ intimately with about four times its measure of the dry and finely-powdered NaCO₃. Then transfer the mixture into the tube and cover it with a layer of the

dry Na₂CO₃.

Now cleanse the inside of the tube perfectly, and proceed to heat the upper layer of Na₂CO₃ strongly, gradually extending the heat to the mixture below, while the upper layer is maintained at a high temperature. Hg-vapour will be evolved, and will condense on the inside of the tube as a silver-white mirror.

Sometimes the globules of metal are so small that they appear as a grey film only, or they are few and scattered and are not easily visible. In either case the minute globules may usually be detected by means of a pocket-lens, or they may be united into larger and visible globules by rubbing the inside of the tube with a glass rod. Even very minute globules may be seen, when they have been flattened by pressing them with a glass rod against the inside of the tube.

Remarks.—The powder is most easily transferred to the tube, by alternately scooping it up with the mouth of the tube, and tapping the tube upon the table so as to shake the powder to the bottom.

Or the powder may be poured upon a narrow strip of notepaper, which has been folded over into a trough longer than the tube; the paper is then pushed into the horizontal tube, the tube is held erect and tapped, and the powder is thus easily transferred to the bottom of the tube without soiling the class.

Both the inside of the tube and the powder must be perfectly dry, else particles will adhere to the inside of the tube and interfere with the detection

of the metallic globules.

The heating must not be commenced until the inside of the tube is perfectly clean. It is cleansed, if necessary, by means of a twisted slip of filter-

paper or with a wooden match.

If any drops of water condense on the inside of the tube during the first stage of the heating, they must be removed by a piece of filter-paper. If, however, proper care has been taken in drying the powder, no moisture will be evolved from it. The escape of steam or gases from the powder is very undesirable, as it is likely to sweep uncondensed mercury vapour out of the tube.

141. Heat a little Solid HgCl₂ in a tube closed at one end; the substance will sublime in white fumes, showing that mercury compounds are volatile.

Bismuth (Bi).

Density 9.8, melting point 268° ('.

142. Bismuth is a brittle, crystalline, white metal with a reddish tinge. It slowly oxidises superficially in the air. The metal is very slightly attacked by hydrochloric acid and by sulphuric acid, but is readily dissolved by nitric acid.

The bismuth salts correspond to the oxide Bi₂O₃ and the bismuthion is trivalent and colourless. Bismuth hydroxide, Bi(OH)₃, is a weak base, hence bismuth salts are partially "hydrolysed" water, forming insoluble basic salts.

For minerals containing Bi refer to par. 1419

REACTIONS OF BISMUTH.

Use Bismuth nitrate, Bi(NO₃)₃.5H₂O, dissolved in dilute HNO₃.

143. H₂S: brownish-black precipitate (Bi₂S₃):--

$$2\text{Bi}(\text{NO}_3)_3 + 3\text{H}_2\text{S} = \text{Bi}_2\text{S}_3 + 6\text{HNO}_3$$

Decant most of the liquid from the precipitate; then shake up the precipitate with the remainder, pour some of the liquid with the precipitate into three test-tubes, and show that the precipitate is insoluble in Am₂S and in KOH, but soluble in strong HNO₃.

- 144. AmOH, or KOH: white precipitate, Bi(OH)₃, insoluble in excess of the reagent. Pour off a small quantity of the liquid and precipitate into a test-tube, add much AmOH and warm: the precipitate will remain undissolved. Filter off the rest of the precipitate, and keep it for the reaction described in paragraph 146.
- 145. Alkaline stannite: add KOH-solution to SnCl₂-solution until the precipitate which forms at first is just redissolved, then add a few drops of the bismuth nitrate solution: a black precipitate of metallic bismuth will be formed.
- 146. H₂O.—Pour a few drops of boiling dilute HCl upon the precipitate which remained in the filter from par. 144, and allow the acid solution to drop from the funnel into a large quantity of distilled water. A milkiness will appear (Note), which is due to the formation of BiOCl:—

$$BiCl_3 + H_2O = BiOCl + 2HCl.$$

, Pour some of the milky liquid into two test-tubes; add to one portion a little strong HCl and warm it, the precipitate will dis-

solve; to the other portion add $H_2\bar{T}$ and warm, the milkiness will remain, differing in this way from that caused by SbOCl (189).

Note.—If Bi is present in small quantity only, this milkiness will often not appear until the liquid has been stirred and then allowed to stand for a few minutes.

A similar precipitate (BiONO₃) may be obtained by allowing the original bismuth nitrate solution to fall drop by drop into much water, but the nitrate is less completely precipitated than the chloride by mixture with excess of water.

147. K₂CrO₄, or K₂Cr₂O₇, added after some strong NaÅ-solution (*Note*), gives a yellow flocculent precipitate of basic bismuth chromate. When KOH is added, the precipitate will not dissolve, differing in this way from PbCrO₄; it will dissolve, however, completely when HCl or HNO₃ is added in excess.

Note.—The addition of NaÅ in the above reaction is necessary in order to replace the nitric acid in the original solution by HÅ, since the precipitate is readily soluble in HNO₃, but is insoluble in HÅ:—NaÅ + HNO₃ = NaNO₃ + HÅ.

148. Blowpipe-test on Charcoal.—Heat a mixture of powdered $Bi(NO_3)_3$ and Na_2CO_3 in a cavity on charcoal in the inner blowpipe-flame.

There will be seen around the cavity an incrustation which is or inge-red while hot and yellow when cold. White globules of Bi will also be seen within the cavity; they are very brittle and will be easily crushed to powder by a blow with the pestle; they are too hard to mark paper (106).

Copper (Cu).

Density 8.9, melting point 1083° C.

149. Copper is a reddish-brown lustrous metal, which is soft, ductile and malleable. In ordinary air it becomes coated slowly, at ordinary temperature, with a green basic carbonate; if it is heated strongly in the air, a superficial coating of black oxide is formed.

Cold hydrochloric acid and sulphuric acid scarcely affect the metal; it is dissolved slowly by hot hydrochloric acid, and rapidly by nitric acid under all conditions. The metal is also rapidly attacked by hot strong sulphuric acid. Ammonia acts upon copper in the presence of air, forming a blue solution.

Copper forms two classes of compounds: the cuprous salts, corresponding to the red cuprous oxide, Cu₂O, contain the monovalent colourless ion, cuprosum; the cupric salts, corresponding to the black cupric oxide, CuO, contain the divalent bluish-green

ion, cupricum.

The cuprous salts are white and insoluble in water, and they readily pass by oxidation in acid or ammoniacal solution into the blue cupric compounds; they are therefore not of common occurrence and are comparatively unimportant: a few distinctive tests only are given for them.

The cupric salts are of common occurrence and of considerable importance, their reactions are therefore described in detail. Their

solutions are blue in colour.

For minerals containing Cu refer to par. 1419.

REACTIONS OF CUPROUS COMPOUNDS.

Use a solution of Cuprous chloride, Cu₂Cl₂, in dilute hydrochloric acid.

- 150. **KOH**: yellow precipitate of $\operatorname{Cu}_2(Off)_2$: when the liquid is boiled the precipitate loses water and changes into red cuprous oxide ($\operatorname{Cu}_2(0)$). The precipitate quickly absorbs oxygen when it is exposed to the air, and is converted into black cupric oxide, CuO .
- 151. H₂S: black precipitate of cuprous sulphide (Cu₂S): soluble in warm dilute nitric acid as blue cupric nitrate, with separation of sulphur.
- 152. $\mathbf{H}_2\mathbf{O}$: when the cuprous salt solution is poured into much water, insoluble white $\mathrm{Cu}_2\mathrm{Cl}_2$ is precipitated, and this gradually turns blue by oxidation when it is exposed to the air.
- 153. The Dry Reactions for cuprous compounds give results similar to those for cupric compounds (161-163).

REACTIONS OF CUPRIC COMPOUNDS.

Use Copper sulphate, CuSO₄.5H₂O.

154. H₂S: brownish-black precipitate (CuS): filter off the precipitate, keeping the funnel carefully covered with a glass plate, since CuS is quickly oxidised by exposure to the air, and the CuS(). thus produced dissolves and runs through in the filtrate.

Place some of the precipitate in five test-tubes and show that it is insoluble in KOH, in Am₂S and in boiling dilute H₂SO₄, but that it is soluble in boiling HNO₃ and in KCN-solution.

155. AmOH, if it is added in a diluted state and in very small

quantity, gives a greenish-blue precipitate.

If more AmOH is added this precipitate dissolves, yielding an intensely blue liquid, which contains a cuprammonium compound, Cu(NH₃)₄SO₄.

" 156. KOH produces a pale blue flocculent precipitate of Cu(OH), which is converted into black CuO when the liquid is heated.

157. KCN-solution: yellow precipitate, CuCy₂, which changes to white Cu₂Cy₂, and this dissolves in excess of the KCy-solution as colourless potassium cuprous cyanide, K₃CuCy₄.

*This solution contains no copper-ion, and is therefore not precipitated by H₂S and by other reagents which precipitate cupric solutions. This property is made use of as a means of separ-

ating copper from cadmium (170,4).

- r58. Potassium ferrocyanide, K_4 FeCy₆: reddish-brown precipitate (Cu₂FeCy₆), insoluble in $H\bar{A}$; the colour of this precipitate is best seen when the reaction is tried in a white porcelain dish. In a very dilute copper solution, only a reddish colour is seen at first when the ferrocyanide solution is added.
- 159, Fe.—Polish a piece of steel or iron wire or sheet, or the blade of a penknife, if necessary, by rubbing it with emery-paper, remove all grease by boiling it in a little dilute KOH-solution, then dip it into CuSO₄-solution, which has been acidified with a few drops of H₂SO₄; the immersed surface will be gradually covered with a red film of Cu.
- 160 Zn and Pt. Place a piece of platinum foil or wire in some CuSO₄-solution, which has been acidified with H₂SO₄ in a porcelain dish or in a watch-glass, then drop a small piece of Zn upon the platinum: a red film of Cu will be deposited upon the platinum.
- 161. Blowpipe test on Charcoal.—If finely-powdered copper sulphate and Na₂CO₃ are mixed in a cavity on charcoal, and the mixture is then heated in the inner blowpipe-flame, red scales or globules of Cu are obtained. These are best seen after they have been washed by levigation (61). The addition of solid KCN facilitates the smelting process, and often enables globules of copper to be obtained.
- 162. A Borax-bead containing Cu is green while hot and blue when cold, after it has been fused in the outer blowpipe-flame.

Fusion in the inner blowpipe-flame causes the bead to become red; it may even become colourless, if very little Cu is present and the bead is long heated in a good reducing flame: the introduction of a minute fragment of Zn or Sn into the bead aids this change.

163. Flame-coloration.—If a loop of platinum wire is dipped into CuSO₄-solution and is then held in the inner blowpipe-flame or in the Bunsén-flame, it imparts a green coloration to the flame.

The flame coloration shows a blue centre after the loop has been moistened with HCl, or if CuCl₂-solution was originally used.

Gadmium (Cd).

Density 8.6, melting point 320.9° C.

164. Cadmium is a white metal, and is fairly malleable and ductile. It is slightly attacked by moist air; and when it is strongly heated, it burns to the brown oxide, (3d).

The metal is readily soluble in the common acids. It forms one series of salts corresponding to the brown oxide, ('dO. The Cd-ion is divalent and colourless.

Cadmium occurs in nature as greenockite, ('dS, and is usually present in small quantity in zinc ble ide.

REACTIONS OF CADMIUM.

Use Cadmium sulphate, 3CdSO₄.8H₂O.

165. H₂S: bright yellow precipitate (CdS). Boil off the H₂S and pour the liquid containing the precipitate into five tubes.

Show with these separate portions that the precipitate is insoluble in Am₂S, in KOH and in KCN-solution, but is soluble in hoiling dilute HNO₃ and in boiling dilute H₂SO₄.

Dilute the H₂SO₄-solution of the precipitate with much water and pass H₂S for a short time; yellow CdS will be reprecipitated.

- 166. AmOH, if it is added in a very dilute condition and in small quantity, gives a white precipitate, Cd(OH)₂; when more AmOH is added, the precipitate will be readily dissolved.
- 167. KOH: white precipitate of Cd(OH)₂, insoluble in excess of KOH.
- 168. KCN: white precipitate of cadmium cyanide, CdCy2, readily soluble in excess as the double cyanide K_2 CdCy4. From this solution H_2 S reprecipitates CdS. (Difference from copper.)
- 169. Blowpipe-test on Charcoal.—If powdered CdS()₄ is mixed with Na₂CO₃ in a cavity on charcoal, and the mixture is then heated in the inner blowpipe-flame, a brown incrustation will be seen on the surface of the charcoal around the cavity.

DETECTION OF THE METALS IN GROUP II.A., WHEN THEY OCCUR TOGETHER.

170. The following Differences are utilised:-

1. The solubility of Bi₂S₃, CuS, and CdS in strong boiling HNO₃, in which HgS and PbS are insoluble and remain as HgS and PbSO₄ respectively after the treatment with the acid.

- 2. The solubility of PbSO₄ in a mixture of HĀ with excess of AmOH, or in AmĀ-solution, HgS being insoluble in AmĀ.
- 3. The solubility of Cu(OH)₂ and Cd(OH)₂, and the insolubility of Bi(OH)₃ in excess of AmOH.
- 4. The solubility of CdS, and the insolubility of CuS in boiling dilute H₂SO₄; or the formation of the double soluble cyanides of these metals with KCN, and the subsequent precipitation of the Cd as CdS by means of H₂S, the copper remaining in solution.
- 171. A Solution which may contain the Metals of Group II.A. is first made acid, if it is not already so, by adding to it HCl. Whether the liquid is clear or not, H₂S is then passed through it for about five minutes, and the precipitated sulphides are filtered off.

The clear filtrate is either mixed with H₂S-water, or it is diluted with a little water and H₂S is passed through it again for a short time, in order to ascertain whether the metals have been completely precipitated from the original solution.

If no further precipitate is produced, the filtrate may be rejected. But in case H₂S causes further precipitation, the gas is passed for some time longer and the liquid is then poured again through the filter. The filtrate is rejected only when H₂S no longer produces any precipitate in it.

The precipitate is then examined by Table II. (1252), commencing at Group II.A. Copper Group (1253), and using only the left-

hand side of the Table.

GROUP II.B.—ARSENIC GROUP.

172. This Group includes As, Sb, Sn.

These metals differ from the metals of Groups III.A., 111.B., 1V., and V., by being precipitated as sulphides by H₂S from a solution which has been acidified with HCl. They differ from metals of Group I. by not being precipitated by HCl.

The sulphides of the metals of this Group differ from those of the metals

in Group II.A. by being soluble in Am2S and in KOH.

Each metal of this Group forms two series of compounds. These compounds resemble each other in many reactions, but they show distinctive differences in certain reactions. The two classes are conveniently distinguished by the terminations -ous and -ic, as in the two classes of mercury and copper salts; thus we speak of arsenious exide and arsenic exide, and of stannous and stannic chloride.

The Reactions of the Rarer Elements of this Group, Au, Pt, Ir, Mo, Se, Te, are given in paragraphs 220-261,

Arsenic (As).

The density is 4.7 in the amorphous state, and to 5.7 when crystalline. Arsenic sublimes without melting at a dull red heat, but it may be fused if it is heated under pressure.

173. Arsenic is a steel-grey metallic-looking substance; in moist air its surface becomes gradually covered with a film of oxide. When arsenic is heated it is converted into yellow vapour, which smells like garlic, and burns with a bright flame emitting white. smoke (As $_4O_6$). Dilute acids have little action on this element, but concentrated acids attack it readily.

The arsenious compounds correspond to the oxide As₄O₆, which is amphoteric in character, yielding with strong acids feeble salts such as AsCl₃, and with strong bases arsenites such as Na₃AsO₃.

Arsenic compounds are converted into arsenicus by boiling them with H.SO₃, with strong HCl, or with some other suitable reducing agent. Arsenicus compounds are oxidised into arsenic compounds when they are nested with HNO₃, or with some other suitable oxidising agent.

For minerals containing As refer to par. 1419.

REACTIONS OF ARSENIC.

Use for an arsenious compound, arsenious oxide (As₄O₆) dissolved in boiling water; and for an arsenic compound, sodium hydrogen arsenate (Na₂HAsO₄.12H₂O) dissolved in water.

Many of the tests described in the following paragraphs give the same ultimate results with both arsenious and arsenio compounds; the reaction, however, usually proceeds more slowly with an arsenio compound, since the first stage consists in its conversion into an arsenious compound.

174. H₂S produces only a yellow coloration in neutral solution of an arsenious compound. But if the solution is acidified with a few drops of HCl, a light yellow flocculent precipitate (As₂S₃) will form immediately even in the cold.

Filter off some of the As₂S₃, and show that it is insoluble in boiling strong HCl.

Pour some of the liquid containing the precipitate into two test-tubes, and add KOH and Am₂S respectively drop by drop; each of these solutions will completely dissolve the As₂S₃ as arsenite and sulpharsenite respectively:—

$$As_2S_3 + 6K_0H = K_3AsO_3 + K_3AsS_3 + 3H_2O.$$

 $As_2S_3 + 3Am_2S = 2Am_3AsS_3.$

The As₂S₃ may be reprecipitated from these solutions by the addition of HCl in excess:—

$$K_3AsO_3 + K_3AsS_3 + 6HCl = As_2S_3 + 6KCl + 3H_2O$$

 $2Am_3AsS_3 + 6HCl = As_2S_3 + 6AmCl + 3H_2S.$

Note.—Since the ammonium sulphide solution used as a reagent is commonly yellow and contains a polysulphide, and the reagent has been added in excess, the As_2S_3 when reprecipitated by HCl will usually be mixed with sulphur: $Am_2S_n + 2HCl = 2AmCl + H_2S + S_{n-1}$.

In cold acidified solutions of arsenic compounds H₂S produces no precipitate. But if strong HCl is added and the solution is boiled, H₂S will precipitate first white S and then yellow As₂S₃, when it is passed for some time into the boiling liquid.

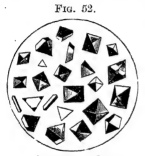
The complete precipitation of arsenic solutions by H₂S is most rapidly secured after they have been converted into the arsenious state. This may be effected by boiling with strong HCl as above, or more rapidly by adding H₂SO₃ to the boiling arsenic solution until it smells persistently of SO₂, and then boiling the liquid until SO₂ is no longer smelt. The arsenious compound thus produced may then be readily and completely precipitated by H₂S in the cold solution.

175. Reinsch's test.—Several narrow strips of sheet copper are cleansed, if necessary, by means of emery-paper, or by dipping them into strong HNO₃. They are then boiled in the arsenious solution, which has been previously acidified by dilute HCl. The surface of the copper will soon be coated with a dark grey film of copper arsenide.

The strips are carefully removed from the liquid, rinsed, and gently pressed between filter-paper; they are then dried by heating them in the steam-oven, or by holding them in the fingers and heating them gently over the flame.

The dry strips are now gradually heated over a very small flame in a warm and dry test-tube. A white sublimate of As₄()₆ will form on the upper part of the tube.

A much more characteristic sublimate will be obtained, if moisture is entirely removed from the copper strips and from the tube, and



Arsenious Oxide Crystals.

if the sublimation is then effected very slowly and while the sides of the tube are hot.

This sublimate will be sparkling and crystalline; and if the crystals are examined under the microscope, they will be seen to be transparent, colourless, regular octahedra and tetrahedra (Fig. 52). The formation of these crystals under the above conditions is a most valuable confirmation of the presence of As.

The sublimate of As O may be dis-

solved in a little boiling water, and the presence of As confirmed by applying tests 174 and 184 to separate parts of this solution.

The presence of As in the original grey deposit upon the Cu should always be confirmed by one of the above methods, since other metals may be deposited as grey films on Cu.

Reinsch's test will serve for the detection of arsenic compounds, after their solutions have been reduced to arsenicus compounds by being mixed with strong HCl or with H₂SO₃-solution and boiled.

Tests which depend upon the Formation of Arsenic hydride (AsH₃).

The following tests (176-180), which depend upon the formation of very poisonous gaseous AsH₃, should be performed in a closet provided with a good fraught: this precaution is necessary in order to guard against any of the gas being inhaled.

176. Hofmann's test.—Pour dilute HCl upon some pure Zn in a small hydrogen-flask, fitted as is shown in Fig. 53.

The hydrogen may be washed, as it escapes, by letting it bubble through some water contained in the second flask, but this is not necessary.

Allow the hydrogen to pass through some AgNO₃-solution con-

tained in a test-tube. If the Zn and acid are free from As, as they should be, no precipitate or colour will be produced in the AgNO₃-solution.

Now pour a few drops only of As_4O_6 -solution down the funnel into the hydrogen flask; a black precipitate of Ag will soon form in the AgNO₃-solution, and H_3AsO_3 will remain dissolved.

This precipitate is produced by the AsH₃, which is formed by the action of the nascent hydrogen upon the As₄O₈:—



HOFMANN'S ARSENIC APPARATUS.

$$As_4O_6 + 12H_2 = 4AsH_3 + 6H_2O.$$

The AsH₃, on passing into the AgNO₃-solution, reacts thus:— $AsH_2 + 6AgNO_3 + 3H_2O = 3Ag_2 + H_3AsO_3 + 6HNO_3.$

As soon as a copious precipitate has been obtained, the testtube is removed.

It will be best to set aside the test-tube with its contents to be examined afterwards, and to proceed at once to use the gas, while it is being briskly evolved, for reactions 177 and 178.

The liquid in the test-tube is filtered; the precipitate is rejected, and several drops of AgNO₃-solution are added to the clear filtrate. Some very dilute solution of ammonia is made by pouring several drops of AmOH-solution into a test-tube, then nearly filling up with distilled water and inverting the tube. When this solution is mixed drop by drop with the filtrate, a light yellow precipitate of silver arsenite will be obtained.

In order to explain the formation of this precipitate, it must be remembered that the solution contains H_3AsO_3 in the presence of $AgNO_3$ and of HNO_3 ; and further that $AgNO_3$ forms a yellow precipitate with H_3AsO_3 if no free acid is present in the liquid (184). The formation of this precipitate is prevented by the HNO_3 ; but when sufficient AmOH is added to neutralise the free HNO_3 which is already present, as well as that which is formed by the action of $AgNO_3$ on H_3AsO_3 , the yellow precipitate will appear:—

 $H_3AsO_3 + 3AgNO_3 + 3AmOH = Ag_3AsO_3 + 3AmNO_3 + 3H_2O_3$

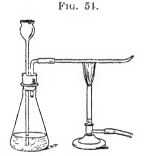
This precipitate (Ag₃AsO₃) is, however, readily soluble in AmOH; hence it is necessary to use very dilute AmOH and to add it gradually in order to avoid excess, since if excess of AmOH is added the precipitate will be prevented from forming.

177. Marsh's test.—Disconnect the flasks (Fig. 53, p. 81), and slip into the rubber joint a tube of glass about four inches (10 cm.) in length, supporting it on the ring of a retort-stand (Fig. 54). Then pour in a little strong, HCl through the funnel-tube.

Caution.—If the experiment is performed as is here directed, the hydrogen will have been already evolved for some time for the previous test (176), and will have expelled all the air from the flask. There will therefore be no risk of an explosion occurring when the gas is lighted, owing to mixture of

air with the hydrogen.

But if the hydrogen is being produced for the first time, or if the flask has been opened or recharged, allow the gas to escape briskly for four or five minutes; and then ascertain that a test-tube, filled with the gas, burns quietly, before proceeding to kindle the gas at the jet. In any case it is well to cover the flask with a cloth before the hydrogen is kindled, in order to guard against any risks from an explosion.



MARSH'S ARSENIC MIRROR.

Now light the hydrogen as it escapes from the jet (Fig. 54). The flame will be seen to differ from that of pure H by its livid blue colour, and by its evolution of white fumes of As₄O₆. If these appearances are not noticed, there is not sufficient arsenic solution in the flask: a few more drops of the arsenic solution must, therefore, be poured in through the funnel-tube.

Press down upon the flame the inside of a porcelain crucible-lid or

of a porcelain crucible or dish. A dusky black film of As will be deposited upon the cool surface. Two porcelain surfaces should be thus stained with As.

Pour upon one of these stains some freshly-made solution of bleaching-powder; the stain will be rapidly dissolved.

Warm the second stain of As with yellow Am₂S. It will dissolve, and the solution will leave *yellow* As₂S₃ when it is slowly evaporated.

178. Decomposition of AsH, by Heat.—While the flame is still burning at the jet, pour a little more strong HCl into the flask (Fig. 54), if necessary, in order to produce a fair-sized flame. Then heat a point near the middle of the tube strongly (Fig. 54). A

black mirror of As will form just beyond the part which is heated by the flame. The flame burning at the jet will meanwhile become colourless, showing that the arsenic has been removed from the hydrogen.

Heat another point in the tube, so as to produce a second mirror; then remove the tube and divide it between the mirrors and confirm the presence of As as follows:—

Dissolve one mirror, when it is cold, by warming it in a testtube with some freshly-made solution of bleaching-powder,

Heat the other mirror gently in a test-tube held obliquely in the flame; if the sides of the tube have been first heated, a crystal-line sublimate of As_4O_6 will be obtained (Fig. 52, page 80). Dissolve this sublimate, when it is cold, in a little hot water and confirm the presence of As by trying reactions 174 and 184 on the solution.

170. HgCl₂-solution is coloured by contact with AsH₃.

Remove the cork from the generating flask (Fig. 53 or 54), and push a plug of cotton-wool into its neck in order to arrest any liquid spirted from the flask. Then cover the mouth of the flask with a small piece of, filter-paper, which has been moistened with HgCl₂-solution; the paper will become coloured, first yellow and then brown. This result is distinctive from SbH₃, which gives only a greyish-brown stain.

If paper moistened with AgNO₃-solution is substituted for the above paper, it will be *blackened* (176): this is also the case with SbH₃.

180. Fleitmann's test.—Boil some pieces of Zn in KOH-solution: H will be slowly evolved, and will be free from odour. If a little As₄O₆-solution is now added and the liquid is again boiled, a garlic odour will be perceptible; the odour is due to the evolution of AsH₃:—

$$\begin{cases} Zn + 2KOH = H_2 + Zn(OK)_2. \\ As_4O_6 + 12H_2 = 4AsH_3 + 6H_2O. \end{cases}$$

If a small piece of filter-paper, which has been moistened with AgNO₃-solution, is placed on the mouth of the tube while the liquid is being boiled, the paper will be stained black by the separation of Ag (176).*

An arsenic compound must be reduced, by being boiled with H₂SO₃, before the above test is applied.

*Sb-compounds do not give the above result; the test therefore serves to distinguish between As and Sb.

REACTIONS OF SOLID ARSENIC COMPOUNDS.

181. Mirror of As by Reduction.—Dry some finely-powdered substance, containing As, thoroughly in the steam oven, and mix it intimately in a mortar with about four times ifs measure of freshly ignited Na₂CO₃, or of a perfectly dried mixture of KCN and Na₂CO₃. Heat this mixture in a little bulb-tube of hard glass (Fig. 55), or in a narrow test-tube about three inches (8 cm.) in length, carefully observing the precautions mentioned in para-



Fig. 55.

ARSENIC MIRROR FROM SOLID ARSENIC COMPOUND.

graph 140. A black mirror of As will form on the cooler part of the tube (B), and a smell of garlic may usually be detected at the mouth of the tube.

If the bulb is cut off and the mirror is gently heated in a testtube, the As will be converted into a white sublimate of As_4O_6 , or into transparent octahedral crystals of the oxide (Fig. 52, page 80).

- 182. Blowpipe-reaction on Charcoal.—Mix a solid substance, containing As, with Na₂CO₃ and KCN, and heat the mixture on charcoal in the inner blowpipe-flame. The flame will be seen to be coloured *livid blue*, unless this tint is overpowered by the yellow sodium coloration. A smell of garlic will be noticed when the charcoal is removed from the flame.
- 183. Heat a little Solid As₄O₆ in a small ignition-tube: it will be readily sublimed. If the As₄O₆ is perfectly dry, and it is sublimed very slowly after the upper part of the tube has been first heated in the flame, the sublimate will sparkle, and when it is examined under the microscope, it will be seen to consist of estabedral crystals (Fig. 52, page 80).

DISTINCTIVE TESTS FOR ARSENIOUS AND ARSENIC COMPOUNDS.

Use As₄O₆ dissolved in boiling water for the arsenious solution, and aqueous solution of sodium arsenate for the arsenic compound.

184. AgNO₃: if several drops are added to the arsenious acid solution, and very dilute AmOH-solution is then added drop by drop, a light yellow precipitate of silver arsenite (Ag₃AsO₂) will form, which is soluble in AmOH and in HNO₃.

With arsenate, a brown precipitate of silver arsenate (Ag₃AsO₄) is produced, which is also soluble in AmOH and in HNO₃.

•185. CuSQ₄, if it is added to the arsenious solution, followed by a gradually increasing quantity of KOH-solution, yields first a yellowish-green precipitate, which then dissolves to a clear blue liquid: this liquid, when it is boiled, deposits orange-red Cu₂O.

With arsenate, the pale green precipitate forms, but it is not soluble in excess of KOH-solution, and it does not deposit Cu₂O

when it is boiled.

186. AmCl, AmOH and MgSO₄: these solutions, added in the order stated, give no precipitate in the arsenious solution.

When they are added to the arsenate solution, a white crystalline precipitate of MgNH₄AsO₄, 6H₂O is produced, and if this is filtered off, and moistened on the filter with AgNO₃-solution, it turns brown (184).

Antimony (Sb).

Density 6.62, melting-point 630° C.

187. Antimony is a brittle, white, lustrous metal. It slowly tarnishes in moist air, and when it is fused and heated in the air it burns to Sb_4O_6 .

Dilute acids scarcely act upon this metal, but concentrated hydrochloric and sulphuric acid convert it respectively into the trichloride and into an unstable sulphate. Hot strong nitric acid converts the metal into the insoluble oxides, Sb₂O₄ and Sb₂O₅.

The antimonious compounds, corresponding to the oxide S b₄O₆ are the important compounds of antimony. This oxide is feebly

basic in character.

The antimonic compounds, corresponding to the oxide $\mathrm{Sb}_2\mathrm{O}_5$, are of little importance to the analyst.

For minerals containing Sb refer to par. 1419.

REACTIONS OF ANTIMONY.

Use Antimony trichloride (SbCl₃) solution, produced by dissolving SbCl₃ or Sb₄O₆ in dilute HCl.

Antimonic and antimonicus compounds give results, which are ultimately identical, in the following reactions (188–195), with the exception of reaction 189. After these reactions, which are common to Sb in both its sets of compounds, two are given which are distinctive of the two classes of compounds (196, 197).

188. H₂S: orange-red precipitate (Sb₂S₃). Pour off the liquid with the precipitate into three tubes, and show that the precipitate

is dissolved when it is warmed with pure KOH or with Am₂S, but is reprecipitated from these solutions when HCl is added in excess (174 and *Note*). Show also that the precipitate is soluble in boiling strong HCl.

189. H₂O.—When SbCl₃-solution is poured into much water, it yields a white precipitate or turbidity (SbO(1): this basic salt is converted into the trioxide when water is added in large amount.

The precipitate is dissolved when HCl is added to the liquid and it is heated; it is also readily soluble in H₂T. The solubility of SbOCl in H₂T distinguishes it from the similarly formed BiOCl precipitate (146).

190. Zn and Pt.—If the acid solution of SbCl₃ is poured upon some platinum-foil in a porcelain dish, and a small piece of Zn is dropped upon the foil, the platinum will be rapidly coated with a black deposit of Sb. If the foil is then rinsed with water and heated in boiling HCl, the stain will remain undissolved; but it will be rapidly removed by heating the foil with HNO₃.

The formation of a slight grey or brown stain on the Pt is nevidence of the presence of Sb. The presence of Sb in the stain may be proved by dissolving it in a little boiling dilute HNO₃, diluting the cold solution, and then passing H₂S and allowing it to stand at rest for some time, when red Sb₂S₃ will settle down.

TESTS WHICH DEPEND UPON THE FORMATION OF ANTIMONY HYDRIDE.

SbH₃ may be generated by pouring Sb-solution into a hydrogen-apparatus, as is directed in paragraphs 176-179, for an arsenic solution. Antimony hydride, SbH₃, is evolved. This gas differs from AsH₃ by being free from smell and from poisonous properties, and also in the respects mentioned in paragraphs 191-193.

The first test in paragraph 179 is also distinctive.

191. If SbH₃ is passed into AgNO₃-solution (176), it produces a black precipitate of SbAg₃:—

$$SbH_3 + 3AgNO_3 = SbAg_3 + 3HNO_3$$
.

Filter off this precipitate, and reject the filtrate which will contain no Sb.

Wash the precipitate of SbAg, four or five times with boiling water upon the filter. Then pour upon it boiling dilute H₂T-solution, which will dissolve the Sb. Receive this liquid in a test-tube, boil it, and pour it once more upon the filter. Now add H(! to the liquid, filter if necessary from AgCl, and pass H₂S into the solution; orange red Sb₂S₃ will be precipitated.

192. A Stain of Sb on Porcelain, which has been produced in the manner already described in paragraph 177, differs from the As-stain by presenting a dead surface and by being insoluble in solution of bleaching-powder. Also when the stain is dissolved by warming it with yellow Am₂S, and the solution is evaporated by a gentle heat, a residue of orange-red Sb₂S₃ is left.

193. The Sb-mirror, obtained as is described in paragraph 178, differs from the As-mirror by being formed much nearer to the flame and on both sides of the heated part of the tube, proving that Sb is less volatile than As. The mirror may further be distinguished by the test which is described in paragraph 191.

The sublimate of Sb₄O₆, which is produced by heating this mirror in the air, differs from the corresponding arsenic sublimate (178) by being almost non-crystalline and insoluble in boiling water. The sublimate may, however, be dissolved in a little boiling HCl, and this acid solution yields an *orange-red* precipitate of Sb₂S₃ when H₂S is passed into it; whereas the solution of the As₄O₆ sublimate yields *yellow* As₂S₃ under similar conditions.

194, Zn and KOH.—If a solution containing Sb is boiled with Zn and KOH, it does not evolve SbH₃; hence the escaping hydrogen does not stain paper moistened with AgNO₃-solution (180). This test serves to distinguish As from Sb in its compounds.

TESTS FOR SOLID COMPOUNDS OF SB.

195. Blowpipe-test on Charcoal.—Mix a little solid substance containing Sb in a cavity on wood-charcoal with Na₂CO₃ and KCN, and heat the mixture in the inner blowpipe-flame.

A white incrustation will be formed on the charcoal; and white globules of metallic Sb will be obtained, which are extremely brittle (106). The flame will also be seen to be coloured pale blue, unless this tint is concealed by the yellow sodium coloration.

When the melted Sb-globules are withdrawn from the flame, they will emit white fumes and become coated with sharp white crystals of $\mathrm{Sb}_4\mathrm{O}_6$.

DISTINCTIVE TESTS FOR ANTIMONIOUS AND ANTIMONIC COMPOUNDS.

Use SbCl₃-solution in HCl, and aqueous potassium antimonate solution respectively.

196. KHO in excess, then AgNO₃: with SbCl₃, a dark coloured precipitate, which when shaken after addition of AmOH leaves black Ag₄O undissolved: with antimonate, a brown precipitate which dissolves entirely on addition of AmOH.

197. HCl in excess, warm, and KI-solution: with SbCl₃ no iodine is set free, the liquid remaining colourless, even after it has been mixed with fresh starch solution: with antimonate, siodine is liberated which colours the liquid brown, and gives a deep blue colour when starch solution is mixed with the cold liquid.

Tin (Sn).

Density varies from 6.6 to 7.3, melting-point 231.9° C.

198. Tin'is a white, lustrous metal, and is both malleable and ductile. At ordinary temperature air has practically no effect upon it, but when the metal is heated above its melting-point in the air it gradually changes into the white dioxide, SnO₂.

Dilute hydrochloric or sulphuric acid slowly dissolves tin with formation of a stannous salt: the hot concentrated acids act more rapidly. Dilute nitric acid forms unstable stannous nitrate, and the concentrated acid forms white hydrated SnO₂.

Tin forms two series of compounds, the stannous and stannic, in which the metal is divalent and tetravalent respectively, and which correspond to the oxides SnO and SnO₂. Both oxides possess feeble basic and acidic properties.

For minerals containing Sn refer to par. 1419.

REACTIONS OF TIN.

Stannous and Stannic compounds.

Use for Stannous salt, SnCl₂.2H₂O dissolved in dilute HCl, and for Stannic salt, SnCl₄ dissolved in dilute HCl.

The reactions 199 and 200 give ultimately the same results with both stannous and stannic compounds. Reactions 201, 202, and 203 serve to distinguish stannous from stannic salts, and to detect stannous salts in the presence of stannic salts.

199. Zinc, when it is immersed in the solution of a tin-salt which has been acidified with HCl, becomes conted with either a spongy mass or beautiful crystals of Sn.

The Zn should be allowed to remain for some time in the solution, which is contained in a small porcelain dish, and the reaction may be hastened by the application of a gentle heat. The deposition of Sn from SnCl₂ takes place more rapidly than from SnCl₄, since SnCl₄ has to be first converted into SnCl₂.

If the Zn rests upon platinum-foil in the solution, the Sn will be deposited in a spongy state on the Zn and will not stain the platinum black. Sn differs in this way from Sb (190).

In general analysis the deposit on the Zn may consist of other substances besides Sn. It should, therefore, be removed by rubbing

and washing the Zn in some water in the porcelain dish; the deposit may then be rinsed into a narrow test-tube, and dissolved in a little boiling strong HCl. This solution will contain SnCl₂, which will yield with HgCl₂-solution a white precipitate of Hg₂Cl₂ (135, 202).

Since this precipitation of $\mathrm{Hg_2Cl_2}$ depends upon the formation of $\mathrm{SnCl_2}$ and its presence in the acid solution, and since $\mathrm{SnCl_2}$ is readily converted into $\mathrm{SnCl_4}$ by oxidation in the acid liquid, precautions must be taken against exposing the acid solution freely to the air. It will usually suffice to dissolve the deposit quickly in a narrow test-tube, and to proceed at once to add the $\mathrm{HgCl_2}$ -solution.

200. Blowpipe-test on Charcoal.—Mix any solid compound, containing Sn, with powdered Na₂CO₃ and KCN, and heat the mixture on charcoal in the inner blowpipe-flame. A slight white incrustation will be obtained, together with white particles of metallic Sn which are with difficulty fused into globules.

The globules of Sn are malleable; they differ from those of Pb by their greater hardness, which prevents them from marking paper. Refer to paragraph 106 for a description of the methods of testing these globules.

DISTINCTIVE REACTIONS FOR STANNOUS AND STANNIC COMPOUNDS.

201. H₂S: dark brown precipitate (SnS) with stannous solution, soluble in boiling strong HCl, also soluble in KOH and when warmed with yellow Am₂S, and reprecipitated as brown SnS from the KOH-solution by HCl, and as yellow SnS₂ from the Am₂S-solution (Note, 174). SnS is insoluble in colourless Am₂S.

With stannic solution, a yellow precipitate (SnS₂) which often forms only when the liquid is heated: soluble both in yellow and in colourless Am₂S and in KOH when heated, and reprecipitated as SnS₂ by HCl from both solutions (Note, 174).

202. HgCl₂: white precipitate of Hg₂Cl₂ with stannous solution, which turns grey when the liquid is boiled if the SnCl₂ is present in excess (135).

No precipitate is produced with stannic solution.

203. AuCl₃ added after a drop of Br-water: with stannous solution a *purple* or *dark brown* precipitate of "purple of Cassius." No precipitate is produced with stannic solution.

DETECTION OF METALS IN GROUP II.B., WHEN THEY OCCUR TOGETHER.

Three methods for the separation and detection of As, Sb and Sn are given. The first, which is described in paragraph 205, is simple and can be rapidly carried out, but the methods of Hofmann (206) and Fresenins (207) are m ore delicate and trustworthy.

- 204. The following Differences are utilised in the first, method:
- 1. The solubility of Sb₂S₃, SnS and SnS₂ in hot strong HCl, in which As₂S₃ is insoluble.
- 2. The action of Zn and Pt in the presence of HCl upon the solution, Sb staining the platinum, and Sn being deposited upon the Zn (190, 199): or the treatment of the H(1-solution with Zn, when the Sn is deposited on the Zn (199), and the Sb is evolved as SbH₃ gas (19f-193).

FIRST METHOD.

205. From a Solution which may contain As, Sb, and Sn. these metals are first precipitated as sulphides by passing H₂S into the acidified solution. The sulphides are then filtered off, and the filtrate is tested by passing H₂S through it again and warming to prove that the precipitation has been complete, this procedure being repeated if necessary. When no further precipitate is produced by H2S the precipitated sulphides are examined in the following manner, based on the above differences (204).

The precipitate produced by H₂S is filtered off, and is then allowed to drain for some time upon the filter in the funnel.

It is best to drain the precipitate still further, either by suction of the filter-pump (44), or by opening out the filter and spreading it upon a pile of three or four cut filter-papers.

The precipitate is then removed to a small porcelain dish, and is heated for some time, nearly to boiling, with strong fuming HCl. The liquid is then cooled and filtered:-

The Residue will consist chiefly of As₂S₃, which is almost insoluble in strong HCl. It may be examined by either I or II below:-

Dry the washed re-nique on the filter of a gentle heat, dam nix it with this or four times as much powdered

The Filtrate may be examined for Sb and Sn by either I. or II. below :--

I. Place a piece of platinum-foil in a porcelain dish and pour the acid filtrate upon it, then place upon the foil a piece of Zn; H will come off with effervescence, and if either at once or after a few minutes a black stain appears upon the platinum this shows :- Presence of Sb.

[The Sb thus deposited on the Pt may be rinsed and then dissolved by heating the foil in a testtube with a little very dilute HNO2; if this solution is cooled and diluted, and HaS is passed through it, an orange-red precipitate will be

formed, confirming the presence of Sb.

KCN and Na₂CO₃, and heat the mixture in a small bulb-tube, removing any drops of water inside the tube by a twisted piece of filter-paper; a black mirror (181):—
Presence of As.

II. Or the As₂S₃ may be dissolved by heating it with a little fuming HNO₃. The excess of HNO₃ is then boiled off and the As is detected as H₃AsO₄ by adding AmCl, excess of AmOH and MgSO₄ (186); or by adding AgNO₃ and then cautiously neutralising with AmOH (184).

Romove the platinum-foil from the dish and place pieces of Zn in the acid liquid; as soon as the bubbles of H cease to be given off, Zn still remaining undissolved, remove the pieces of Zn, rubbing and rinsing any dark deposit back into the dish: let this deposit settle, decant the liquid, and heat the solid deposit with strong HCl for several minutes in a test-tube, dilute with a little water, filter if necessary, and add a few drops of HgCl₂-solution; a white or grey precipitate (135, 202) indicates:—Presence of Sn. The original solution may then be examined to find whether the Sn was present as stannous or stannic salt (201-203.)

II. The acid filtrate is poured into a little hydrogen flask (Fig. 54, p. 82), in which H has been coming off briskly for about five minutes, being produced by the action of a little strong HCl upon some pieces of pure Zn. The H is lighted at the jet, and the inside of a small porcelain dish or crucible lid is pressed down upon the flame; a black stain, which is not dissolved by solution of bleaching-powder, shows:—Presence of Sb.

The residue in the flask is tested for Sn, as is directed in the latter part of I. (above).

SECOND METHOD.

206. Hofmann's method depends upon the different results obtained by the action of Zn upon the dissolved chlorides of the metals in the presence of HCl. The Sn is deposited upon the Zn, and AsH₃ and SbH₃ gases are evolved. These gases are allowed to react upon silver nitrate solution, when the As passes into solution as H₃AsO₃ and the Sb is precipitated as SbAg₃.

The sulphides, which have been precipitated as is directed in paragraph 205, are warmed with a little strong HCl, and a small crystal of KClO₃ is occasionally dropped in until solution is complete. The warming of the liquid is then continued until no smell of Cl is evolved.

This solution of the metals is then poured into a suitable flask, in which H is being generated by the action of HCl upon pure Zn (Fig. 53, p. 81). Sn, Au and Pt, if present, will be deposited upon the Zn, while As and Sb will be evolved as gaseous AsH₃ and SbH

3. These two gases are passed into AgNO₃-solution, and yield H

3AsO₃ in solution and Ag

3Sb as a black precipitate.

The AgNO3-solution, containing the black precipitate, is filtered:-

Precipitate: wash this well upon the filter, and pour upon it some boiling H₂T-solution. Boil this liquid again and pour it several times on the filter. Now add HCl and pass H₂S: a red coloration or precipitate indicates:—Presence of Sb.

Filtrate: add to this some very dilute AmOH, drop by drop, with constant mixing by stirring or shaking; a pale yellow floculent, precipitate indicates:—Presence of As.

The Deposit of Sn is removed from the Zn in the flask, and is washed. It is then dissolved by boiling it with a little strong HCl in a narrow test-tube: any residue being examined by paragraph 1305, if An and Pt have to be tested for. The solution in HCl is at once diluted and is tested for SnCl₂ by the addition of HgCl₂-solution: a white precipitate, possibly turning grey (135, 202), shows:—Presence of Sn.

A further test is made, if necessary, upon the original solution in order to ascertain whether the tin was present as stannous or as stannic salt (201-203).

THIRD METHOD.

207. Fresenius' method depends upon the fact that fusion with Na₂CO₄ and NaNO₃ converts As₂S₃ into soluble Na₂HAsO₄, Sb₂S₃ into insoluble Na₂H₂Sb₂O₇, and SnS and SnS₂ into insoluble Sn or SnO₂.

The Sulphides, which have been precipitated as is directed in paragraph 205, are dried on the filter, and the precipitate is well mixed with equal weights of powdered Na₂CO₃ and NaNO₃.

If the quantity of the precipitate is very small, the part of the paper which contains the precipitate is cut up into small pieces, and these are treated as

is directed for the precipitate.

This mixture is dropped in small quantities at a time into some NaNO₃, which is fused in a small porcelain crucible over a Bunsen-flame. When the powder has all been added, the crucible is heated sufficiently to keep the mixture melted for a few minutes, and the liquid mass is then poured into a small dry porcelain dish.

As soon as the substance has cooled, cold water is poured upon it in the dish, and water is also poured upon the residue which adheres to the inside of the crucible. The solid substance is allowed to soak in the water in both vessels for some time with occasional stirring, crushing it by pressure with a pestle if it does not easily fall to pieces; the liquid is then filtered:—

The Residue is washed on the filter with a mixture of equal measures of water and alcohol, in which Na₂H₂Sb₂O₇ is insoluble; the washings are rejected.

The precipitate is then rinsed into a small porcelain dish, using as little water as possible; a few drops of strong HCl are added and the dish is heated, more HCl being added, if necessary, to make the liquid acid to litmus-paper.

Whether the residue has dissolved or not, place in the liquid a piece of clean platinum-foil, and drop upon the foil a piece of pure Zn. Remove the foil after a few seconds: a black stain shows: Presence of Sb. The presence of Sb may be confirmed by dissolving it off the foil with a little hot dilute HNO₃, cooling and passing H₂S, when a red precipitate of Sb₂S₃ will form.

When the Zn ceases to be acted upon, Sn if present will remain as a spongy deposit. This is to be washed by decantation, and then dissolved by boiling with HCl in a test-tube: HgCl₂ is added immediately to the solution: a white precipitate (202) shows:— Presence of Sn. The original solution is then tested, if necessary, to find whether Sn was present in the stannous or stannic condition.

Any residue insoluble in HCl should be examined for Au and Pt (1305).

Filtrate: HNO3 is added until the solution iн after acid being boiled; then AgNO_a-solution poured in as long as it causes any precipitate, and very dilute AmOH is added little by little (Note); brown precipitate, readily dissolving in **OXCORS** of AmOH shows :--(184), Presence of As.

Note. — Instead of mixing dilute AmOH with the liquid, AmOH may be carefully poured upon the top of the liquid; a brown colour is then seen at the surface of contact.

REACTIONS OF THE RARER ELEMENTS IN GROUP II.

208. The Rarer Elements of this Group are divided into two classes: those belonging to Group II.A. are Pd, Os, Rh, Ru; those belonging to Group II.B. are Au, Pt, Mo, Se, Te, Ir.

GROUP II.A.—COPPER GROUP.

209. In this Group are included Pd, Os (Rh, Ru); these metals are precipitated as sulphides by $\rm H_2S$ from acid solutions, and the sulphides are insoluble in $\rm Am_2S$ and in solution of alkaline hydroxide.

Palladium (Pd).—Use solution of Palladious chloride, PdCl2.

- 210. Pd occurs as a metal in native platinum, also in gold and in silver: it has a density of 11.4. Palladious solutions are reddish-brown, and become yellow on dilution: the addition of water precipitates from these solutions a basic salt, unless sufficient free acid is present to prevent the precipitation.
- Palladic compounds are also known, such as PdCl₄; they give the same reactions as palladious compounds, since they readily change into the palladious state.
- **211.** $\mathbf{H}_2\mathbf{S}$: black precipitate in neutral, alkaline, and acid solutions; this precipitate is insoluble in $\mathrm{Am}_2\mathbf{S}$, but soluble in boiling HCl and in aqua regia.
- 212. AmOH: flesh-coloured precipitate Pd (NH₃)₂"Cl₂; soluble in excess of AmOH to a colourless liquid, from which HCl precipitates the isomeric yellow crystalline palladammonium chloride.
- 213. $HgCy_2$: yellowish-white gelatinous precipitate ($PdCy_2$), slightly soluble in HCl, easily soluble in AmOH and in KCN-solution. This is a very characteristic reaction.
 - 214. KI: black precipitate (PdI₂); a very characteristic reaction.
- 215. Dry test: all Pd-compounds are decomposed by ignition, leaving the metal as a residue.

Osmium (Os).—Use solution of K₂OsCl₆. Density of the metal 22.48.

- 216. H₂S: brownish-black precipitate of OsS, insoluble in Am₂S.
- 217. KOH, AmOH or K2CO3: reddish-brown precipitate of Os(OH) 4.
- 218. Tannic acid or Alcohol, when it is added to the solution previously acidified with HCl, reduces the osmic chloride to blue osmious chloride, OsCl₂.
- 219. Formation of OsO₄: after acidifying a little of the solution with dilute HNO₃, distil it and receive the distillate in KOH-solution. The alkaline solution will be coloured yellow by the formation of potassium osmiate. Acidify this solution, and note the penetrating odour of the liberated OsO₄. Also add a little Na₂S₂O₃-solution to a portion of the distillate: when this is warmed a brown precipitate of osmium sulphide will be produced.

GROUP II.B .- ARSENIC GROUP.

220. In this Group are included Au, Pt, Mo, Se, Te, (Ir).

Gold (Au).-Use solution of Gold chloride.

- 221. Metallic gold is yellow; it has a donsity of 19-3, and melts at \$4064°; it is the most ductile metal known. Gold is unaffected by air under all conditions. It is insoluble in all the common liquids and acids, but is dissolved by chloring- or bromine-water, and by aqua regia. Two unstable, weakly basic oxides are known, Au₂O and Au₂O₃. The light yellow solution of gold in aqua regia contains both AuCl₃ and HAuCl₄, and is known as gold chloride solution: it is used for analytical and photographic purposes.
- 222. H₂S: in φld solutions black Au₂S₂ is precipitated, soluble in yellow Am₂S; in hot solutions brown metallic gold is precipitated.
- 223. SnCl₂ containing a little SnCl₄: purple or dark brown precipitate of "Purple of Cassius," which is best seen when the test is made in a white porcelain dish. This precipitate consists of very finely divided gold mixed with hydrated stannous oxide.
- 224. FeSO₄-solution produces, either at once, or when the solution is heated, a very finely-divided precipitate of Au. The particles of the precipitate cannot be seen, but they usually cause the liquid to appear bluish by transmitted light, and always copper-red and turbid by reflected light: —

$$AuCl_3 + 3FeSO_4 = Au + Fe_2 (SO_4)_3 + FeCl_3$$

 $225.\ H_2O_2$: in alkaline solution gold is immediately precipitated in a finely-divided brownish-black condition: the liquid only appears reddish in colour if the gold is in very small quantity, and no precipitate is seen. This reaction is very delicate.

Platinum (Pt).—Use solution of Platinic chloride, or chlorplatinic acid, H₂PtCl₆.

226. Platinum has a density of 21.5, and melts at 1755° C. The metal is greyish-white in mass, but is known also as a black finely-divided powder "platinum black." Platinum is not acted on by air under any conditions but forms a brittle carbide when it is strongly heated in contact with soot or hydrocarbon gases. No ordinary acids affect it, but it is slowly acted on by chlorine- or bromine-water, and is readily dissolved by aqua regia, yielding a reddish-yellow solution which contains H₂PtCl₆, chlorplatinic acid. This solution is used for analytical purposes under the name of platinic chloride solution.

Platinum forms two unstable feebly basic oxides, PtO and PtO2.

- 227. H₂S: black precipitate (PtS₂), which forms slowly in the cold, but more rapidly when the liquid is warmed. The precipitate is soluble in yellow ... Am₂S and in aqua regia.
 - 228. AmCl, added to a strong solution, produces on standing for some time, or more quickly on being stirred, a yellow precipitate (Am₂PtCl₃), which is soluble in much hot water.
 - 220 KCF yellow crystalline precipitate of K2PtCl6: the addition of a

- 230. Formic acid precipitates from neutral boiling solutions all the platinum as a black powder: an acid solution must be previously neutralised by sodium hydroxide solution.
- **231.** SnCl₂ gives a reddish-brown colour (H_2PtCl_4) in the solution acidified with $HCl := H_2PtCl_6 + SnCl_2 = H_2PtCl_4 + SnCl_4$.

The Method of Separating and Detecting Au and Pt is fully given in paragraph 1305.

Iridium (Ir).—Use solution of Na₂IrCl₆.

- 232. H_2S : the dark red colour of the solution is discharged, sulphur is deposited, and eventually a brown precipitate of $Ir_2S_0^{\dagger}$ is deposited: the precipitate readily dissolves in Am_2S .
- 233. KOH: the colour of the cold solution changes to green; and when the solution is warmed its colour first changes to reddish, and finally to azure blue:— $2 \text{IrCl}_4 + 2 \text{NaOH} = 2 \text{IrCl}_3 + \text{NaCl} + \text{NaOCl} + \text{H}_2\text{O}$.
 - 234. KCl: brownish-black precipitate of K2IrCl6, insoluble in alcohol.
- 235. Reducing reagents, $SnCl_{2}$ or $FeSO_4$ -solution, change the colour of the solution to green by reducing the $IrCl_4$ to trichloride; or they precipitate the metal as a black powder, as is the case with Zn.

Molybdate (MoO₄).—Use Am₂MoO₄-solution.

- 236. Mo occurs as molybdate: also as sulphide, which may be readily converted into MoO₃ by heating it in the air with HNO₃. Unignited MoO₃ dissolves in acids; ignited MoO₃ is insoluble in acids, but is easily soluble in alkali and in ammonium hydroxide solution.
- 237. HCl, HNO₃, or H_2SO_4 , if it is added in small quantity to an aqueous solution of a molybdate, yields a white precipitate of H_2MoO_4 which is readily soluble in excess of the acid.
- 238. H_2S , added in very small quantity to the acidified solution, gives a blue liquid; if H_2S is added in larger quantity, a brown precipitate (MoS₃) forms; the precipitation becomes complete only when the solution is heated and H_2S is passed for some time. This precipitate is soluble in solution of alkaline sulphide or of alkaline hydroxide; MoS_3 is reprecipitated from this solution when acid is added in excess.
- 239. Zn, or SnCl₂, if it is added to the solution of a molybdate in HCl, colours it brown, green, or blue, according to its state of concentration
- 240. KCNS, if it is added to a solution acidified with HCl, gives no coloration; but if a piece of Zn is afterwards dropped into the liquid, a beautiful crimson colour is produced: when this liquid is shaken with ether, the colour is taken up by the ether.
- 241. Na₂HPO₄, if it is added in very small quantity to the solution of molybdate acidified with HNO₃, gives a yellow precipitate of ammonium phospho-molybdate when the liquid is gently warmed; this precipitate is readily soluble in excess of alkaline hydroxide solution.

- 242. Borax-bead : outer flame, yellow; inner flame, dark brown.
- 243. Microcosmic-bead: outer and inner flames, green.

Selenium (Se).—Use PbSe, K₂SeO₃, and K₂SeO₄.

244. Se occurs as selenide of Pb, Fe, Cu, Ag: also in small quantities in pyrites, and when this pyrites is burnt for the manufacture of sulphuric acid the Se is deposited in the leaden chambers. Selenium forms selenides, selenites and selenates, which are analogous to the corresponding sulphur compounds.

I. Selenide (Se.)-5Jse PbSe.

245. When a Selenide is heated in an open tube it evolves a smell of decaying horse-radish, and produces a grey or red sublimate of Se: the smell is very characteristic.

II. Selenite (SeO₃).—Use K₂SeO₃ (251).

- **246.** H_2S gives in a cold acid solution a yellow precipitate, and in a hot acid solution a reddish-yellow precipitate: $-H_2SeO_3 + 2H_2S Se + S_2 + 3H_2O$. This precipitate is soluble in Am_2S .
- 247. BaCl $_2$ gives in a neutral solution a white precipitate (BaSeO $_3$) soluble in HCl and in HNO $_3$.
- 248. SnCl₂ or H₂SO₃ gives in the presence of free HCl a red, or in warm solutions a grey precipitate of Sc.
- 249. Cu in a hot solution acidified with HCl becomes covered with a black film, and the liquid, on standing with the Cu for some time, becomes coloured red by Se.
 - 250. CuSO4: green crystalline precipitate: selenates give no precipitate.

III. Selenate (SeO_4) .—Use K_2SeO_4 .

- 251. HCl produces no change in the cold; but when the liquid is boiled, Cl is given off and the selenate is reduced to selenite, to which the above tests (246-250) may be applied:— $K_2SeO_4 + 2HCl = K_2SeO_3 + Cl_2 + H_2O$.
- 252. BaCl₂: a white precipitate (BaSeO₄); insoluble in cold HCl; dissolved by boiling with HCl, with evolution of Cl:—BaSeO₄ + 4HCl = $\rm H_2SeO_3 + BaCl_2 + Cl_2 + H_2O$. This behaviour with boiling HCl serves to distinguish and to separate the BaSeO₄ precipitate from the similar BaSO₄-precipitate.

IV. Selenium in any form of Combination.

- 253. When it is heated on Charcoal in the Innar Elowpipe-flame a solid selenium compound evolves a smell of rotten horse-radish.
- 254. When it is heated on Charcoal with Na₂CO₃ in the Inner Blowpine-flame a selenium compound yields a fused mass, which when cooled and moistened on Ag yields a black stain Ag₂Se, and on addition of HCl evolves H₂Se, a badly smelling gas.

Tellurium (Te).—Use PbTe, K2TeO3, K2TeO4, as is directed below.

255. Te commonly occurs united with the metals Au, Ag, Bi, Cu, Fe.

Telluride (Te).—Use Lead telluride, PbTe.

256. When a Telluride is heated in an open glass tube, it gives white amos and a sublimate; the sublimate differs from that given by Sb, by being fusible before the blowpipe.

II. Tellurite (TeO₃)".—Use K₂TeO₃ (260).

257. $\mathbf{H}_2\mathbf{O}$: on dilution with water, tellurous acid is precipitated from the acid solution.

258. $\mathbf{H}_2\mathbf{S}$: in acid solutions a brown precipitate (TeS₂), easily soluble in Am_2S .

259. $\mathbf{H}_2\mathbf{SO}_3$, \mathbf{SnCl}_2 , or \mathbf{Zn} precipitates black Te from a solution which is not strongly acid.

III. Tellurate (TeO₄)".—Use K₂TeO₄.

260. HCl produces no change in the cold solution of a tellurate; but when the liquid is boiled, Cl is evolved and the tellurate is reduced to tellurite, the solution then gives the above reactions (257–259).

IV. Tellurium in any form of Combination.

261. If it is fused with Na₂CO₃ on Charcoal in the Inner Blowpipe-flame, a tellurium compound gives sodium telluride, a solution of which gives a black stain on Ag. When this solution is acidified it deposits black Te, and evolves H₂Te.

For the Detection of the Rarer Elements in Group II., see the second column in Table, paragraph 1323.

GROUP III.—IRON AND ZINC GROUPS.

262. The Group includes Al, Fe, Cr, Zn, Mn, Ni, Co, together with the rarer metals U, In, Be, Ti, V, Zr, Ce, Ta, Nb, La, Di, Yt, E, Th. It is conveniently subdivided into Groups III.A. and III.B. The reactions of the rarer metals of this group are given in paragraphs 326-397.

GROUP III.A.—IRON GROUP.

263. The Group includes Al, Fe, Cr.

The members of this Group are not precipitated by the group-reagents for Groups I and II and they differ from those of Groups III.B., IV., and V. by being precipitated from their solutions when AmOH is added in excess after AmCl. This Group is also precipitated by Am₂S, or by H₂S added in excess to the amminiscal solution.

Group III.A. further differs from Group III.B. by being completely precipitated by adding BaCO₃ shaken up with water. This reagent affords the most perfect means of separating these two Sub-groups from one another, unless the metals are present in solution as sulphates when both groups are precipitated.

The members of Group III.A. show no characteristic flame-colorations; but, with the exception of Al, they impart a characteristic colour to the borax-bead.

Aluminium (Al).

Density 2.58, melting point 658.7° C.

264. Aluminium is a white, lustrous metal, and is very slightly acted upon by moist air; it is malleable and ductile, but not very tenacious.

The metal dissolves slowly in cold hydrochloric acid, but rapidly when heated: nitric acid has scarcely any action upon it. Cold sulphuric acid scarcely affects the metal, but the hot concentrated acid dissolves it. Alkaline hydroxide solutions rapidly dissolve the metal with evolution of hydrogen and formation of aluminates:— $Al_2 + 6KOH = 3H_2 + 2K_3AlO_3$.

Aluminium forms only one series of salts, and the trivalent ion is colourless and feebly basic.

For minerals containing Al refer to par. 1419.

REACTIONS OF ALUMINIUM.

Use Ammonium alum, AlAm(SO₄)₂.12H₂O.

265. AmOH: colourless flocculent precipitate Al(OH)_B, which is often transparent and invisible until it has been coagulated by boiling the liquid: it is soluble in HCl and in HĀ, and is slightly soluble in AmOH in the absence of AmCl.

266. KOH added in very small quantity: colourless flocculent precipitate, Al(OH)₂, soluble in excess of KOH:—

$$AJ(OH)_3 + 3KOH = K_3AIO_3 + 3H_2O.$$

Add dilute HCl gradually to part of the KOH-solution of the precipitate until the liquid is neutral, the Al(OH)₈ will be reprecipitated. If more acid is added the precipitate will dissolve, but will reappear when AmOH is added in slight excess.

From another portion of the KOH-solution of the precipitate the Al(OH), may be reprecipitated by the addition of sufficient AmCl-solution.

267 Ammonium sulphide, Am,S: the same precipitate of M(OH), H.S gas being evolved or remaining dissolved in the

 $\Delta H_{3}(SO_{4})_{2} + 3Am_{3}S + 6H_{3}O = 2Al(OH)_{2} + 3Am_{3}SO_{4} + 3H_{3}S.$

268. Blowpipe-test.—Heat some solid alum on charcoal in the outer blowpipe-flame; then moisten it, after cooling, with $Co(NO_3)_2$ -solution and again heat it in the outer flame; a *fine office miss* will be obtained.

Iron (Fe).

Density 7.8, melting-point 1500° C.

269. Metallic iron readily oxidises or "rusts" in moist air, and is eventually converted into hydrated ferric oxide.

When the metal is heated strongly in air, or in oxygen, it is converted into the black oxide, Fe₃O₄.

Iron readily dissolves in acids: hydrochloric and sulphuric acids yield the corresponding ferrous salts, but nitric acid produces a ferric salt and evolves reddish-brown gases.

Commercial iron is never pure. It usually contains small quantities of carbon, sulphur, phosphorus, silicon and manganese, and frequently traces of other substances. When commercial iron is dissolved in HCl or in H₂SO₄ the hydrogen which escapes is accompanied by small quantities of gases, such as hydrocarbons, hydrogen sulphide, and hydrogen phosphide, and these impart to the gas an unpleasant odour. There is always a residue left after the action of the acid is complete, which consists mainly of carbon.

For minerals, containing Fe refer to par. 1419.

270. Two classes of iron-compounds are known, corresponding to the oxides FeO and Fe_2O_3 . They differ in appearance and properties and behave differently with reagents. They are known as *ferrous* and *ferric* compounds respectively. Ferric salts are usually yellow or reddish-yellow. Ferrous salts are usually pale green if they are in solution, or are combined with water (hydrated), and are white if they are perfectly free from water or "anhydrous."

The metal in ferrous compounds is divalent and is often distinctively termed ferrosum; in ferric compounds the metal is trivalent and may be termed ferricum.

271. It is usual to ascertain in which state of combination the iron exists, and some distinctive tests are therefore described below.

The whole of the reactions of ferrous compounds are not given, because Fe is always separated and detected in the course of analysis as a ferric compound, and ferrous compounds are subsequently tested for by trying special tests upon the original substance.

Ferrous compounds are readily converted into ferric compounds by heating them for a short time with an oxidising agent, such as a little strong HNO₃, or HCl and a crystal of KClO₃, or Br-water.

Both classes of compounds give the same results in the blowpipe-reactions (275, 276).

REACTIONS OF IRON.

Use FeCl₃ and FeSO₄.7H₂O.

272. AmOH, or KOH: with ferric salt the Fe is completely precipitated as a reddish-brown flocculent precipitate, Fe(OH)₃, which is insoluble in KOH, and soluble in HCl.

With ferrous salt the Fe is only partially precipitated as a dingy-green precipitate, which ultimately turns to brown Fe(OH)₃ on exposure to the air.

273. Am₂S: a black precipitate (FeS) is obtained with both ferrous and ferric salts, but it is mixed with white sulphur in the case of ferric salts:—

$$2\text{FeCl}_3 + 3\text{Am}_2\text{S} = 2\text{FeS} + \text{S} + 6\text{AmCl}.$$

FeS is soluble in boiling acids, but is insoluble in KOII.

In very dilute iron-solution a green colour only is produced at first by Am₂S, but black FeS separates after the liquid has stood for some time or when it is boiled.

274. H₂S, either passed as gas into the solution, or added as H₂S-water, gives a black precipitate (FeS) when the iron-solution is subsequently mixed with excess of alkaline hydroxide.

In neutral or acid solution of ferric salt, sulphur is precipitated and the ferric salt is converted into ferrous salt; the colour of the solution therefore changes from yellow to pale green: this change of colour is visible in a strong solution, after it has been boiled and filtered:— $2FeCl_3 + H_2S = 2FeCl_2 + 2HCl + S$.

- 275. Heated with Solid Sodium carbonate (Na₂CO₃) and Potassium cyanide (KCN) on Charcoal in the inner blowpipe-flame, a solid iron compound leaves a grey powder of metallic iron (60). This may be separated by levigation (61), and shown to be mignetic by being attracted when it is touched under water with the end of a magnet, or with the point of a magnetised knife-blade.
- 276. A Borax-bead containing Fe is reddish-brown while hot and yellow when cold, after it has been fused in the outer blowpipe-flame: after fusion in the inner-flame it is olive-green.

DISTINCTIVE TESTS FOR FERROUS AND FERRIC SALTS.

Use acidified solutions of FeCl₃ and FeSO₄.

- 277. KOH or AmOH gives distinctive precipitates (272).
- 278. Potassium ferrocyanide, K.FeCy.: with ferric solution

dark blue precipitate, "prussian blue," insoluble in HCl and turned brown by KOH-solution.

With ferrous solution a *light blue* precipitate, which becomes dark blue by oxidation in the air or when it is treated with HNO₃ or Br-water.

279. Potassium ferricyanide, K₃FeCy₆: in freshly made solution (*Note*) darkens the colour of a ferric solution, but produces no precipitate, as is seen when the liquid is diluted.

With ferrous solution a dark blue precipitate, "Turnbull's blue," is produced, which is insoluble in HCl. A dark bluish-grey colour

only is produced in very dilute ferrous solution.

Note.—This solution must be made immediately before it is used, by dissolving a small piece of solid potassium ferricyanide, about the size of a pin's head, in a few drops of water.

280. Potassium thiocyanate, KCNS: with ferric solution a blood-red coloration, but no precipitate: the presence of HNO₃ or of HĀ hinders the reaction. The colour is immediately destroyed when a few drops of the red liquid are mixed with HgCl₂-solution.

With ferrous solution no coloration is produced by KCNS unless ferric salt is present in small quantity.

Chromium (Cr).

Density 6.92, melting-point 1505° C.

281. Metallic chromium is very hard and has the appearance of cast iron. When it is heated in the air to a high temperature it is converted into Cr₂O₃. The metal is readily soluble in hydrochloric acid and in sulphuric acid, but is practically insoluble in nitric acid.

There are two well-known series of chromium compounds: the chromic salts, which correspond to the basic oxide Cr_2O_3 , and in which Cr is trivalent; and the chromates which contain the acidic oxide CrO_3 .

The chromic salts, which are usually purple in colour, give the reactions which are described in paragraphs 282-287.

•The chromates, which are usually yellow or reddish in colour, give the reactions which are described in paragraphs 639-644.

The chromium salts pass by oxidation into chromates in reactions 285 and 286, where the oxidation is caused by PbO₂ and by KNO₃ respectively.

Chromates, on the other hand, pass by reduction into green chromic compounds; examples of this change will be found under the tests for chromates (639-644).

For minerals containing Cr refer to par. 1419.

REACTIONS OF CHROMIUM.

Use Chrome alum, CrK(SO₄)₂.12H₂().

- 282. AmOH: pale bluish-green or purple precipitate, $Cr(OH)_3$. If AmOH is added in large quantity and the liquid is heated, some of the precipitate will dissolve, producing a beautiful violet-red solution. The colour of the liquid is best seen after the precipitate has been removed from it by filtration. From this red solution the $Cr(OH)_3$ is reprecipitated when the liquid is boiled for several minutes in a porcelain dish.
- 283. KOH, if it is added in small quantity, gives the same precipitate, $Cr(OH)_3$. If more cold KOH is added, the precipitate is entirely dissolved to a green fluid. When this liquid is diluted with water and then boiled for several minutes, the $(r(OH)_3)$ is represented and the liquid becomes colourless. When Am(1 is edded to the KOH-solution, the $(r(OH)_3)$ is also reprecipitated.
- 284. Am₂S also precipitates Cr(OH₃), H₂S gas being given off or remaining dissolved. See the equation in paragraph 267.
- 285. Lead peroxide (PbO₂), or Red lead, may be added to the green liquid which has been obtained by adding KOH in excess to the chromium solution (283). When this liquid is boiled with the lead oxide, its colour changes to yellow; a yellow precipitate (PbCrO₄) may then be thrown down from this yellow solution by adding to it HĀ in excess. This serves as a confirmatory test for the presence of Cr.
- 286. A Solid substance containing Cr, if it is heated with fused Na₂CO₃ and potassium nitrate (KNO₃) on platinum or porcelain, as is described in Exp. 25, par. 56, yields, when it is cold, a bright yellow mass containing sodium chromate (Na₂CrO₄).

When this cold product is dissolved in boiling water, a yellow liquid is obtained; if this solution is acidified with HĀ its colour reddens, and if it is then boiled for a few minutes in order to drive off CO₂, it gives with solution of lead acetate (PbĀ₂) a yellow precipitate of PbCrO₄.

287. A Borax-bead containing Cr is green after it has been fused either in the outer or in the inner blowpipe-flame.

DETECTION OF THE METALS IN GROUP III.A., WHEN THEY OCCUR.

288. The following Differences are utilised:-

1. The solubility of Al(OH)₃ in boiling KOH, in which Fe(OH)₃ and Cr(OH)₃ are insoluble.

- 2. The conversion of Cr(OH)₃ by fusion with Na₂CO₃ and KNO₃ into soluble Na₂CrO₄: Fe(OH)₃, when it is thus treated, remaining as Fe₂O₃ which is insoluble in water.
- 3. Cr is then detected by the yellow colour of the aqueous solution of the resulting Na_2CrO_4 , and by the formation of a yellow precipitate when this solution is acidified with $H\bar{A}$, and $Pb\bar{A}_2$ -solution is added to it.
- 4. Fe is found by the blood-red coloration which is obtained, when KCNS is added to the solution in HCl of the Fe₂O₃ resulting from (2).
- 289. A Solution which may contain Al, Fe, Cr, is first examined for ferrous and ferric salts in the following manner:—
- (a) To a few drops of the liquid, acidified with HCl, add a little K₄FeCy₆-solution: if a blue precipitate is produced, Fe is present: proceed to (b); if no blue precipitate forms, proceed to (c). portion of the liquid is rejected.
- (b) Two other small portions of the acidified solution are then tested; one with freshly made solution of K₃FeCy₆ (Note, 279), which, by giving a dark blue precipitate, would show the presence of Fe⁻⁻; the other with KCNS-solution, which may cause a blood-red colour, proving the presence of Fe⁻⁻. These portions of the liquid are rejected. Proceed to (c).
- (c) The rest of the original solution is boiled, after a few drops of strong HNO₃ (271) have been added if ferrous salt is present. AmCl in volume equal to one quarter that of the solution is then added; then AmOH is gradually added, with constant stirring, until the liquid is just alkaline and smells faintly of NH₃. The liquid is now boiled and filtered, and the precipitate is examined by Table III.A. (1255), all the Notes appended to the Table being disregarded.

GROUP III.B.—ZINC GROUP.

*290. This Group includes Zn, Mn, Ni, Co.

Compounds of these metals are not precipitated by the Group-reagents for Groups I., II., and III.A., and differ from those of metals of Groups IV. and V. by being precipitated from their neutral or alkaline solutions by yellow ammonium sulphide (Note), and from their alkaline solutions by H_2S . Mn is, however, gradually oxidised and precipitated, if its solution is mixed with AmCl and excess of AmOH and is then exposed to the air.

The compounds of the metals of this Group show no characteristic flame-

colorations; but, with the exception of those of Zn, they give characteristic colours to a borax-bead.

Finely-powdered barium carbonate (BaCO₃) does not precipitate the metals of Group III.B. from their solutions, as it does the metals of Group III.A., unless they are present as sulphates.

Note.—The yellow ammonium sulphide, which is repeatedly referred to in the tests for Ni and Co, is the reagent which is ordinarily found in the laboratory. It is a polysulphide of varying composition and may be represented by the general formula Am₂S_n. The monosulphide, Am₂S, and the ammonium hydrogen sulphide, AmHS, are colourless; they may be used as reagen'ts unless yellow Am₂S or Am₂S_n is specified.

Zinc (Zn).

Density from 6.9 to 7.2, melting-point 419.4° C., boiling-point 916' C.

291. Zinc is a white metal which is somewhat ductile and malleable. In moist air it is slowly oxidised at ordinary temperature, and is eventually covered superficially with a film of basic carbonate. When the metal is heated to a high temperature in the air, it burns with a greenish light and is converted into ZnO.

The pure metal is practically insoluble in all acids, but ordinary commercial zinc is readily dissolved by acids and by boiling caustic alkali solution.

The zinc compounds correspond to the basic oxide ZnO, and the soluble salts contain a colourless, divalent ion.

For minerals containing Zn refer to par. 1419.

REACTIONS OF ZINC.

Use Zinc sulphate, ZnSO_{4.7}H₂O.

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292. Am₂S: white precipitate (ZnS). This precipitate ofter appears yellow, owing to the presence of excess of yellow ammonium polysulphide (Am₂S_n) in the liquid. The true colour is evident, as soon as the precipitate, produced by Am₂S_n, has been separated from the liquid by settling or by filtration.

The colour and the solubility of this precipitate are more satisfactorily ascertained when the precipitate is obtained as in the next paragraph (*Note*, 293).

. 293. H₂S: white precipitate (ZnS). Zn is only partly precipitated from the neutral solution by H₂S, since H₂SO₄, in which ZnS is soluble, is formed during the reaction:—

$$ZnSO_4 + H_2S = ZnS + H_2SO_3$$

If, however, NaOH or AmOH has been added in sufficient excess to the zinc solution, it will form a clear solution (294, 295); and since free H.SO. cannot exist in this alkaline liquid, the Zn may be entirely precipitated from it as sulphide by H.S.

The addition of a sufficient quantity of solution of sodium acetate $(Na\bar{A})$ to the zinc solution also enables H_2S to effect complete precipitation of the Zn as sulphide, since acetic acid $(H\bar{A})$, in which 2nS is insoluble, replaces the H_2SO_4 in the solution:—

$$ZnSO_4 + H_2S + 2Na\bar{A} = ZnS + Na_2SO_4 + 2H\bar{A}.$$

Add to some ZnSO₄-solution AmCl, then AmOH until it is alkaline, and then H₂S. White ZnS will be precipitated. Add to separate portions of this liquid HCl, HĀ, and Am₂S respectively; ZnS will be found to be soluble in HCl, but insoluble in HĀ and in Am₂S.

Note.—The tests which show the colours and solubility of the sulphides of the metals of this Group are best tried on the precipitate which is obtained by the addition of AmCl, AmOH in excess and H₂S to the solution of the metal, unless freshly prepared colourless Am₂S can be obtained; since the yellow polysulphide solution causes sulphur to precipitate together with the metallic sulphide:—

$$ZnSO_4 + Am_2S_2 = ZnS + S + Am_2SO_4$$

and the sulphur, thus precipitated, is not dissolved by the solvents of the metallic sulphide. Sulphur is also precipitated on the addition of acid to the liquid containing the precipitate together with an excess of Am₂S_n, and this further complicates the results:—

$$Am_2S_n + 2HCl = 2 AmCl + H_2S + S_{n-1}$$

294. KOH, added in small quantity, gives a white precipitate, $Zn(OH)_2$. The addition of more KOH causes this precipitate to dissolve, owing to the formation of a soluble alkaline zincate:— $Zn(OH)_2 + 2KOH = K_2ZnO_2 + 2H_2O$. When much water is added to this solution and the liquid is boiled, the $Zn(OH)_2$ is reprecipitated. The Zn may also be precipitated from the KOH-solution as ZnS by the addition of H_2S .

295. AmOH, if added in small quantity, yields a precipitate of $Zn(OH)_2$, which readily dissolves in excess: the formation of this precipitate is prevented by the previous addition of AmCl.

296. Blowpipe-test on Charcoal.—Mix some finely-powdered ZnSO₄.7H₂O and Na₂CO₃ in a small cavity on wood-charcoal, and heat the mixture in the inner blowpipe-flame.

An incrustation will be obtained on the charcoal, which is yellow while hot and white when cold. (Note.) This deposit cannot be driven away by the outer blowpipe-flame, but it is easily removed when it is heated in the inner flame.

Moisten the cold incrustation with cobalt nitrate solution and heat it strongly in the outer blowpipe-flame; it will become green.

Note.—The above changes of colour are more distinctly seen when a little zinc oxide (ZnO) is heated in an ignition tube, or in the outer blowpipe flame on charcoal. The oxide will be yellow while it is hot and white when it has cooled. If the oxide is moistened on charcoal with cobalt nitrate

solution, and is then reheated in the outer blowpipe-flame, it will become green.

Manganese (Mn).

Density 8.0, melting-point 1225° C.

297. Manganese is a brittle grey metal, which is slowly oxidised superficially in the air, and decomposes water with evolution of hydrogen. It is attacked by dilute acids, even by acetic acid.

It forms six oxides, MnO, Mn₂O₃, Mn₃O₄, MnO₂, (MnO₃) and

 Mn_2O_7 .

The most important compounds are the manganous compounds, which correspond to the basic oxide Mn() (298-301); and the permanganates, which correspond to the acidic oxide Mn₂()₇ (604-607).

The former compounds contain a divalent pale-pink positive ion, and the latter a monovalent purple-red negative ion.

For minerals containing Mn refer to par. 1419.

REACTIONS OF MANGANESE.

Use Manganous chloride (MnCl₂), or Manganous sulphate (MnSO₄).

Manganous salts are usually of a pale pink colour, alkaline manganates are green, permanganates are purple.

·298. Am₂S: flesh-coloured or pale pink precipitate (MnS). The true colour of this precipitate is only seen after it has been separated by filtration, if the ammonium sulphide was yellow.

The colour of the moist filtered precipitate changes by oxidation to dark brown, when it is allowed to stand in contact with the air.

For the solubility of this precipitate, refer to the next paragraph (Note, 293).

299. H₂S precipitates pale pink MnS, partially from neutral solutions and entirely from alkaline solutions, but it produces no precipitate in the presence of HCl or of HĀ.

Add to some MnCl₂-solution AmCl, AmOH in excess, and then H₂S; and show with separate portions of the precipitate that MnS is soluble in HCl and in HĀ, but is insoluble in Am₂S.

- 300. KOH: white precipitate, Mn(OH), insoluble in excess, and girckly turning brown by oxidation in the ar. This change of colour is most rapid after the precipitate has been filtered off, then it is then exposed more fully to the air.
- AmOH gives the same precipitate, Mn(OH). If sufficient AmOH is first added, however, AmOH produces no immediate pre-

cipitate; but if this alkaline liquid is exposed to the air, it gradually turns brown by oxidation, and the Mn is ultimately precipitated as a brown hydroxide.

302. Blowpipe-test by Oxidation.—When any solid substance containing Mn is heated upon platinum foil in the tip of the Bunsen-flame or of the outer blowpipe-flame, in contact with a fused mixture of Na₂CO₃ and KNO₃, the product yields a bluishgreen mass when it cools.

The mixture of the substance with the alkaline salts should be heated by allowing the tip of the flame to touch the *under surface* of the foil, and the flame must not be allowed to come into contact

with the surface of the mixture (Exp. 25, par. 56).

The test may also be made by fusing a mixture of Na₂CO₃ and KNO₃ into a bead in a loop of platinum wire, then taking up *very little* of the manganese compound upon the surface of the moistened bead, and fusing it into the bead by heating it in the extreme tip of the outer blowpipe-flame.

This reaction for the presence of manganese is very delicate, owing to the intensity of the colour of the manganate which is produced: therefore very little of the manganese compound should be used else the colour of the fused mass may appear almost black, and the result is then not characteristic.

- 303. Blowpipe-test on Charcoal.—When any manganese compound s heated with fused Na_2CO_3 on charcoal in the inner blowpipe-flame it yields a grey magnetic powder of Mn (60, 61).
- 304. A Borax-bead which contains very little Mn, will be riolet-red while it is hot and amethyst-red when cool, after it has been fused in the outer blowpipe-flame. When the bead is heated n the inner blowpipe-flame it becomes colourless.

This reaction is very delicate, and yields a result with even ninute quantities of manganese compounds. Larger quantities of manganese render the bead opaque, after it has been heated in the outer blowpipe-flame.

Nickel (Ni).

Density 8-8, melting-point 1452° C.

305. Nickel is a hard, white magnetic metal, which does not alter in the air. It is dissolved slowly by hydrochloric acid and by ulphuric acid, and rapidly by nitric acid. The metal forms me series of compounds corresponding to the basic oxide NiO, and its soluble salts contain a green divalent ion.

For minerals containing Ni refer to par. 1419.

REACTIONS OF NICKEL.

Use Nickel sulphate, NiSO₄.7H₃O.

Hydrated nickel salts are usually bright green in colour: the anhydrous salts are yellow.

306. Am S: black precipitate (NiS).

Add more yellow ammonium polysulphide, warm and filter. The filtrate will be coloured brown by NiS, which has been dissolved

by the excess of the polysulphide.

Pour this dark-coloured filtrate into a porcelain dish and boil it for some time, adding distilled water if necessary in order to prevent the liquid from evaporating to dryness. The dissolved NiS will be reprecipitated, and when it has been filtered off, the filtrate will be colourless.

If HĀ is added in excess to the dark-coloured filtrate, the dissolved NiS will also be reprecipitated.

For the solubility of NiS, refer to the next paragraph (Note, 293).

307. H₂S: black precipitate (NiS) in a neutral solution, or in a solution which contains no free acid except HĀ (293); HCl and other acids prevent the precipitation.

Add to some NiSO₄-solution AmCl, then AmOH in excess and then H₂S; and show with the precipitate thus produced that NiS is insoluble in cold HCl and in HĀ, but is dissolved when it is heated with HCl and a crystal of potassium chlorate (KClO₃).

- 308. KOH: light green precipitate, Ni(OH)2, insoluble in excess.
- 309. AmOH, added in very small quantity, gives a bluishgreen precipitate of basic salt. This precipitate is soluble in excess of AmOH to a violet-blue liquid: it is also soluble in AmCl.

If AmCl is first added to the NiSO₄-solution, the addition of AmOH causes no precipitate.

- 310. Solution of Potassium cyanide (KCN), if it is freshly-prepared by dissolving a small piece of solid KCN in water, and is added in small quantity to the nickel solution, gives a yellowish-green precipitate (NiCy₂). The further addition of the KCN-solution dissolves this precipitate as K₂NiCy₄, but the precipitate is reproduced when sufficient HCl is added to the solution.
- 311. Precipitation as Ni(OH)₃ from K₂NiCy₄.—Acidify state NSO₄-solution with several drops of HA; then add freshly-made KCN-solution gradually, while stirring or shaking the nickel solution, until the precipitate, which forms at first, is just rediscolved.

Boil this solution for a short time: then cool it, and divide it into two parts.

The addition of HCl in excess to one part will produce a precipi-

tate of NiCy, which often appears only after a time.

A black precipitate, $Ni(OH)_3$, will be formed on the addition to the other part, either of sodium hypochlorite solution (NaClO), or of potassium hypobromite (KBrO):— $2K_2NiCy_4 + KBrO + 5H_2O = 2Ni(OH)_3 + KBr + 4KCN + 4HCN$. The hypobromite is conveniently prepared by adding to the nickel cyanide solution first KOH in excess, and then bromine water until the liquid remains yellow. The black precipitate forms more rapidly if the liquid is warmed.

- 312. Blowpipe-test on Charcoal.—When a mixture of finely-powdered nickel sulphate and $\rm Na_2CO_3$ is heated on charcoal in the inner blowpipe flame, a grey powder of Ni will remain, which may be separated by levigation and shown to be attracted by the magnet (60, 61).
- 313. A Borax-bead containing Ni is violet or reddish-brown immediately after it has been fused in the outer blowpipe-flame, and it becomes pale yellow on cooling.

If the bead is heated for some time in the inner blowpipe-flame,

it becomes black and opaque.

Cobalt (Co).

Density 8.9, melting point 1490° C.

314. Cobalt is a white, hard, magnetic metal, which closely resembles nickel in its properties. The compounds of cobalt correspond to the basic oxide CoO, and its soluble salts contain a pink divalent ion.

For minerals containing Co refer to par. 1419.

REACTIONS OF COBALT.

Use Cobalt nitrate, Co(NO₃)₂.6H₂O.

Hydrated or dissolved cobalt salts are usually reddish-pink in colour. The solid anhydrous salts are usually blue or purple. Cobalt iodide is green.

- 315. Am₂S: black precipitate (CoS). If yellow ammonium polysulphide is added, and the liquid is then boiled and filtered, the filtrate is coloured yellow by Am₂S_n; it is not dark coloured, since CoS, unlike NiS, is insoluble in yellow Am₂S_n. For the solubility of CoS refer to the next paragraph (Note, 293).
- 316. H₂S: black precipitate (CoS), forming only in alkaline solutions, or in solutions which contain no free acid except HĀ (293). The presence of HCl prevents the precipitation.

Add AmCl, then AmOH in excess, and then H₂S to some cobalt nitrate solution, and pour into two separate portions of this liquid HCl and HĀ respectively; the precipitate will not dissolve. To the portion containing HCl add a crystal of KClO₃ and heat; the precipitate will dissolve readily.

- 317. KOH: blue precipitate of a basic cobalt salt; if the liquid is warmed, or allowed to stand, the basic salt is decomposed, yielding the pink hydroxide Co(OH)₂, and this eventually becomes brown by oxidation into cobaltic hydroxide, Co(OH)₃.
- 318. AmOH: bluish-green precipitate, having the same properties as that produced by KOH. The precipitate is soluble in excess of AmOH and in AmOl, and is therefore not formed if AmOl is added to the cobalt-solution before AmOH, or if AmOH is added in excess.
- 319. Freshly-prepared KCN-solution, if it is added in small quantity, gives a reddish-brown precipitate (CoCy₂). When more KCN-solution is added slowly and while shaking the liquid, the precipitate dissolves as K₄CoCy₆, but the original precipitate reappears when HCl is added to the solution.
- 320. Cobalticyanide reaction.—Acidify some cobalt nitrate solution with a few drops of HĀ; then add freshly-made KCN-solution slowly, until the precipitate which forms at first is just redissolved.

Boil this liquid, then cool and divide it into two parts. Add to one of these portions HCl; and to the other either NaClOsolution, or excess of NaOH-solution followed by sufficient Brwater to render the liquid permanently yellow; no precipitate will appear in either case, even when the liquid is warmed (311).

This difference in behaviour of solutions of Ni and of Co, which have been boiled with excess of KCN, is due to the fact that NiCy₃ forms with KCN a compound (K_2 NiCy₄) which is easily decomposed: whereas CoCy₂ forms with excess of KCN a similar compound (K_4 CoCy₆), but this, in the presence of atmospheric oxygen, changes into the very stable compound K_3 CoCy₆, potassium cobalticyanide, which resists decomposition by acids and by alkaline oxidising solutions:—

$$2K_4CoCy_6 + 2H_2O + O = 2K_3CoCy_6 + 2KOH + H_2O.$$

321. Potassium nitrite (KNO₂), when it is added to solution of cobalinitrate acidified with acetic acid, gives a golden yellow crystalline precipitate of potassium cobaltinitrite, $K_8\text{Co(NO_2)}_6$. If the solution is dilute, the precipitate only appears after a time, but its formation is hastened by rubbing the sides of the vessel with a glass rod: (Difference from nickel.)

Blowpipe-test on Charceal.—When a solid cobalt compound is mixed with Na_iCO₃, and the mixture is fused on charceal in the inner blow-pipe flame, it yields a grey magnetic powder of Co (6e, 61).

323. A Borax-bead, which has been dipped into a strong solution of cobalt nitrate, assumes a *fine blue* colour when it is fused in either the inner or the outer blowpipe-flame. This colour is produced even by minute quantities of cobalt compounds.

DETECTION OF THE METALS IN GROUP III.B., WHEN THEY OCCUR TOGETHER.

324. The following Differences are made use of :-

1. The solubility of NiS in yellow ammonium polysulphide in which ZnS, MnS and CoS are insoluble.

NiS may then be reprecipitated by boiling this solution or by the addition of HĀ, and the presence of Ni may be confirmed by heating the NiS in a borax-bead (313).

- 2. The solubility of ZnS and MnS in cold dilute HCl, in which NiS and CoS are almost insoluble.
- 3. The solubility of $Zn(OH)_2$ in cold KOH, in which $Mn(OH)_2$ is insoluble.

White ZnS may then be precipitated from this alkaline solution by the addition of H₂S: and the Mn(OH)₂ may be heated in a fused mixture of Na₂CO₃ and KNO₃, when it will yield a bluishgreen mass (302).

4. The difference in behaviour of the solution of NiCy₂ and of CoCy₂ in excess of KCN-solution, when the liquid is warmed with NaClO or with NaBrO (311, 320).

Another method for separating Ni and Co depends upon the fact that while Co is precipitated as $\mathrm{Co}(\mathrm{OH})_3$ from its solution by BaCO_3 in the presence of Br, Ni is not precipitated by this means and remains in solution.

5. The marked difference in colour of the borax-bead containing Ni from that containing Co is a further means of detecting these metals.

Mn may be separated from Zn, Ni, and Co by passing H₂S into the neutral solution which has been mixed with sufficient NaÅ (293); Mn alone remains in solution (299), and the other metals are precipitated as sulphides.

325. A Solution is Examined for Zn, Mn, Co, Ni, by adding AmCl, then AmOH in excess, and then yellow Am₂S_n until the liquid smells of the sulphide after it has been well stirred or shaken.

The liquid is then heated, and a few drops of it are poured upon a filter. If the filtrate is colourless, sufficient Am₂S_n has not been added; more Am₂S_n must then be added, and the liquid must be again heated and filtered, these processes being repeated until a coloured filtrate is obtained.

As soon as the filtrate is brown or yellow in colour, proving that Am₂S_n has been added in excess, the whole of the liquid is heated and filtered, and the precipitate is examined by Table III.B. (1257).

If the filtrate is yellow it is rejected, if it is brown it must be examined for Ni according to paragraph 1248.

REACTIONS OF THE RARER ELEMENTS IN GROUP III.

326. The rarer metals which are precipitated in the Am₂S group are *Be, *Ti, Ta, Nb [*Ce, *La, *Zr, *Th, *Y, Yt, Er, *Di], *In, *U, *V (Tl). The metals whose symbols are included in square brackets are often called the "metals of the rare earths."

The reactions of those metals whose symbols are marked with an asterisk are described below.

The following motals are also precipitated as hydroxides or acids by AmOH:—Be(OH₂), H_2 TiO₃, H_3 TaO₄, H_3 NbO₄, $Ce(OH)_3$, $La(OH)_3$, $Zr(OH)_4$, Th(OH)₄, Yb(OH)₃, Yt(OH)₃, $Er(OH)_3$, Di(OH)₃; and if Am₂S is subsequently added, U, In, V and Tl are precipitated as $(VO_2)S$, InS, V_2S_3 and Tl₂S respectively.

BaCO3, precipitated, or in finest powder, suspended in water is a usoful

reagent in this group.

TI is often partially precipitated as chloride in Group I., and its reactions are given under that Group. V is not precipitated unless acid is added in excess, after Am₂S has been added.

BERYLLIUM (Be).—Use Beryllium sulphate, BeSO_{4.4}H₂O.

- 327. Be occurs as silicate in phenacite, and as silicate with aluminium silicate in beryl and in emerald. Be-salts in aqueous solution have an acid reaction and possess a sweet, astringent taste. The metal has a density of 2·1.
- 328. Am₂S, AmOH; KOH, or NaOH precipitates the floculent hydroxide $Be(OH_2)$ which resembles $Al(OH)_3$ in appearance and in being soluble in KOH:— $Be(OH_2) + 2KOH = Be(OK)_2 + 2H_2O$. It differs from $Al(OH)_3$ in being precipitated from its solution in KOH by dilution and long boiling, and by the freshly-precipitated hydroxide being dissolved when it is boiled for some time with AmCl-solution.
- 329. Alkaline carbonates precipitate a carbonate, which is soluble in excess of the reagent; the carbonate is reprecipitated when this solution is diluted and boiled for some time; the solution and reprecipitation are most easily effected when Am₂CO₃ is used as the reagent. [Difference from Al(OH)₃).]
- 330; BaCO₃, suspended as fine powder in water, precipitates Be-solutions completely in the cold as Be(OH)₂.
- 331. H₂C₂O₄ and Alkaline oxalates produce no precipitate. (Difference from the metals of the rare-earths.)

332. Moistened with Cobalt nitrate solution and heated on charcoal in the outer blowpipe-flame, a grey mass is obtained. (Difference from Al.)

TITANIUM (Ti).—Use Titanium sulphate, $Ti(SO_4)_2$, or a solution of titanium hydroxide in hydrochloric acid.

- 333. Ti occurs as TiO₂, together with traces of Fe, Mm and Cr, in rutile, anatase, and brookite. TiO₂ occurs also in combination with Fe in titaniferous iron ore.
- 334. Ignited TiO_2 is insoluble in water and in most acids: it is easily soluble in HF, less readily in boiling strong H_2SO_4 : it forms $Ti(SO_4)_2$, which is soluble in cold water, after it has been heated with fused KHSO₄. TiO_2 differs from SiO_2 by not being volatilised when it is heated in a platinum dish with HF and strong H_2SO_4 .
- 335. By Dilution and long Boiling, white flocculent hydrated TiO₂ is precipitated from solution of Ti in H₂SO₄ or in HCl, and from the cold aqueous solution prepared after treatment with fused KHSO₄ (334): the precipitate is metatitanic acid, H₂TiO₃: it cannot be filtered off until either an acid or AmCl has been added. This hydrolytic reaction is used to effect the separation of Ti from Al, Fe, Cr, and other substances.
- 336. AmOH, KOH, NaOH, Am₂S, or BaCO₃, white flocculent precipitate, insoluble in excess; if the precipitation and washing have been carried out in the cold, the precipitate is orthotitanic acid, H₄TiO₄, and dissolves in HCl and in dilute H₂SO₄. If the precipitation is made in hot solutions, almost insoluble metatitanic acid, H₂TiO₃, is precipitated.
- 337. Zn or Sn gives in strong and acid solutions a blue coloration; if the solution is dilute, a rose-red coloration is produced. The change of colour is due to the formation by reduction of coloured compounds, in which Ti is trivalent. The solution is preferably made acid with HCl.
- 338. Na₂S₂O₃ precipitates Ti solution entirely when the liquid is boiled. (Difference from Fe.)
- 339. Microcosmic-bead: in the outer flame yellow while hot, colourless when cold: in the inner flame yellow while hot, violet when cold.

The production of these colours is promoted by introducing a minute fragment of Sn into the fused bead. The addition of a very small quantity of FeSO₄ causes the bead to become blood-red when it is heated in the inner blowpipe-flame.

CERIUM (Ce).—Use Cerous nitrate, Ce(NO₃)₃.

340. Ce occurs as silicate in cerite, orthite and gadolinite; also as phosphate in monazite sand.

Ce forms two oxides only, Ce_2O_3 and CeO_2 , and both of these are basic. The cerous salts are white and are readily soluble in water. The ceric salts are orange-red, are insoluble or sparingly soluble in water, and are readily reduced to cerous salts by ordinary reducing agents: they are unimportant for ordinary purposes

- 341. AmOH or Am₂S: white precipitate of ('e(OH)₃, insoluble in excess, but readily soluble in acids. The precipitate becomes yellow on exposure to air by formation of (e(OH)₄. In the presence of H₂T the above reagents give no precipitate. (Distinction from Yt.)
- 342. KOH: white precipitate of Co(OH)3 turning yellow on exposure to the air.
- 343. $\mathbf{H}_2\mathbf{C}_2\mathbf{O}_4$ or $\mathbf{Am}_2\mathbf{C}_2\mathbf{O}_4$: white precipitate of cerous oxalate, insoluble in excess (Distinction from Zr), also insoluble in dilute mineral acids. White insoluble (CeO₂ is formed when this precipitate is ignited.
 - 344. BaCO3: complete precipitation takes place on standing.
- 345. KClO, KBrO, or Am₂S₂O₈: a yellow or red precipitate on boiling, due to oxidation into ceric compounds.
- 346. H₂O₂, with sufficient AmOH to make the solution faintly alkaline, gives a dark orange precipitate of hydrated CeO₃. This is a most delicate reaction of cerium.
- 347. K_2SO_4 produces a white crystalline precipitate of the double salt, $Ce_23SO_4.3K_2SO_4$ (Distinction from Be and Al), which is insoluble in a saturated solution of K_2SO_4 . (Difference from Yt and Er.)
- 348. Borax-bead: outer flame, red while hot, colourless when cold; inner flame, colourless, but if CeO₂ is used a yellow opaque bead is obtained.

LANTHANUM (La).—Use Lanthanum nitrate, La(NO3)a.

- 349. La occurs in several minerals, but chiefly in gadolinite and monazite. It forms one oxide, La_2O_3 , which is basic and forms colourless salts.
- 350. AmOH, or Am₂S: a white basic salt is precipitated which is difficult to filter.
- 351. KOH: white precipitate of La(OH)3, insoluble in excess; unchanged by oxidising agents. (Distinction from Ce.)
- 352. $\mathbf{H_2C_2O_4}$, or $\mathbf{Am_2C_2O_4}$: white precipitate, insoluble in excess or in dilute mineral acids.
 - 353. Am₂CO₃: white precipitate, insoluble in excess. (Distinction from Ce.)
- 354. K₂SO₄: white crystalline precipitate of La₂(SO₄)₃.3K₂SO₄, insoluble in concentrated K₂SO₄-solution,

La-compounds give no characteristic dry reactions.

ZIRCONIUM (Zr).—Use Zirconium sulphate, Zr(SO4)2.

355. Zr occurs mainly as zircon, ZrO_2 .Si O_2 . Zirconsum forms two oxides, ZrO_2 and Zr_2O_5 ; the former is the more important.

In sulphate may be obtained by fusing zircon with excess of Na₂CO₃, when softwar silicate and Na₂ZrO₄ are formed. On treatment with water the softwar silicate is dissolved, and the sodium zirconate is hydrolysed, leaving use their washed and dissolved in dilute sulphuric acid. The $Zr(OH)_4$ is their washed and dissolved in dilute sulphuric acid.

- 356. AmOH, KOH, or Am₂S: white gelatinous precipitate of Zr(OH)₄, insoluble in excess, but readily soluble in dilute acids if it has been precipitated in the cold. If it is precipitated from a hot solution, it is difficult to readily soluble in strong acids.
- 357. Am₂CO₃: white precipitate of basic carbonate, soluble in excess, but reprecipitated when this liquid is boiled.
- 358. $\mathbf{H}_2\mathbf{C}_2\mathbf{O}_4$: white precipitate of zirconium oxalate, readily soluble in excess and in $\mathrm{Am}_2\mathbf{C}_2\mathbf{O}_4$ -solution. (Distinction from Th and Co.)
 - 359. Na₂S₂O₃: complete precipitation as Zr(OH)₄ mixed with sulphur.
- 360. Turmeric-paper, after being moistened with the Zr-solution acidified with HCl, becomes reddened when it is dried. (Difference from Th.)
- 361. Dry reaction.— ZrO_2 becomes very luminous when it is heated in, the blowpipe-flame.

THORIUM (Th).—Use Thorium sulphate, Th(SO₄)₂.

- 362. Th occurs as thorite, ThSiO₄, also mixed with other rare earths in gadolinite and monazite. Thorium salts correspond to the only oxide ThO₂.
- 363. AmOH, KOH, or Am₂S: white precipitate of Th(OH)₄, insoluble in excess, but readily soluble in dilute acids.
- 364. Am_2CO_3 : white precipitate of basic carbonate, readily soluble in excess, but reprecipitated from this solution when it is heated to 50° C.
- 365. K_2CO_3 or Na_2CO_3 : white precipitate of the carbonate, readily soluble in excess; this solution of the precipitate becomes turbid when it is boiled, but becomes clear again as it cools.
 - 366. BaCO₃ completely precipitates thorium salts in the cold.
- 367. $\mathbf{H}_2\mathbf{C}_2\mathbf{O}_4$: white precipitate, $\mathrm{Th}(\mathbf{C}_2\mathbf{O}_4)_2$, insoluble in excess and in dilute acids, but soluble in excess of hot $\mathrm{Am}_2\mathbf{C}_2\mathbf{O}_4$ -solution. From this hot solution of the double oxalate, HCl precipitates all the Th as oxalate. (Distinction from Zr.)
- 368. K_2SO_4 : white precipitate of $Th(SO_4)_2.2K_2SO_4.2H_2O$, difficultly soluble in water and insoluble in concentrated K_2SO_4 -solution. (Difference from Yt.)
- 369. HF, or KF: white gelatinous precipitate, which becomes heavy and granular. (Difference from Al, Be, Ti, U and metals of the rare earths except Y.)

Y'TTRIUM (Yt).—Use Yttrium nitrate, Yt(NO3)3.

- 370. Yttrium is an important constituent of gadolinite, it is also present in cerite, thorite and monazite.
- 371. AmOH, KOH, or Am_2S : white precipitate of $Yt(OH)_3$, insoluble in excess
- 372. Am₂CO₃, K₂CO₃ or Na₂CO₃ · white precipitate of yttrium carbonate, readily soluble in excess; after this solution of the precipitate has stood for a time an insoluble double salt precipitates.

- 373. $\mathbf{H}_2\mathbf{C}_2\mathbf{O}_4$: white precipitate of the oxalate, insoluble in excess, difficultly soluble in HCl and slightly soluble in $\mathrm{Am}_2\mathbf{C}_2\mathbf{O}_4$ -solution.
- 374. K₂SO₄: no precipitate, since a double salt is formed which is soluble n excess of K₂SO₄-solution. (Difference from Ce, La, Zr, Th.)
- 375. HF: white amorphous precipitate, which becomes pulverulent when t is warmed. (Difference from Al, Be, U, Ti and metals of the rare earths except Th.)
 - Indium (In).—Use Indium sulphate, $\ln_2(SO_4)_3$.
- 376. Indium occurs in zinc-blende and in wolfram: it is a white nalleable metal of 7.42 density.
- 377. Alkaline hydroxides precipitate white, voluminous In(OH)₃, which is insoluble in excess. Indium solutions are also precipitated by alkaline arbonates, by Na₂HPO₄, by boiling their neutral solutions with excess of NaA, by BaCO₃, and by alkaline oxalates.
- 378. H_2S gives no precipitate in strong and acidified solutions; in dilute and feebly acid solutions a little sulphide separates; in a solution, containing to free acid but $H\bar{A}$, the metal is entirely precipitated as yellow sulphide, n_2S_3 . This precipitate is insoluble in cold Am_2S , but dissolves when the Am_2S is boiled: white indium sulphide separates as the hot solution cools.
- 379. Am₂S, added after $H_2\overline{T}$ and excess of AmOH, gives a white precipitate of In_2S_3 , which is soluble in Am₂S; the white precipitate becomes yellow when it is treated with HA.
- 380. Flame-Coloration: bluish-violet. The spectrum shows a very characteristic blue line (Fig. 51, p. 50); when indium chloride is introduced nto the flame this line is brilliant, but it very rapidly disappears.
- 381. Heated on charcoal with Na₂CO₃: white malleable globules of in are obtained.

URANIUM (U).—Use Uranyl nitrate, (UO2)(NO3)2.

382. U occurs in nature principally as pitchblende, U_3O_3 ; also as ur.mite, hydrated uranium calcium phosphate; and as chalcolite, hydrated uranium sopper phosphate.

The metal is heavy, having a density of 18-68.

- Although salts of uranium corresponding to the oxide UO₂ are known, they are extremely unstable and change on exposure to air into uranyl (UO₂) salts. The oxide UO₂ acts as a divalent basic radicle in the stable uranyl salts, and the reactions described below are those of the uranyl compounds. All uranyl compounds are yellow or green in colour.
- 383. AmOH, KOH, or NaOH: yellow precipitate of uranate, Am₂U₂O₇, insoluble in excess.
- 384. Am₂S gives in neutral solutions a dingy yellow or brown precipitate of transit sulphide (UO₂S), which differs from ZnS, MnS, and FeS by being soluble in Am₂CQ₃. The precipitated sulphide settles slowly, unless AmCl is added; it is soluble in soids, even in HA; when it is hered with Am₂S in excess, it is changed into black uranous oxide (UO) and sulphur.

. H2S produces no precipitate in acid solutions of uraryl salts.

- 385. Am₂CO₃, KHCO₃, or NaHCO₃: yellow precipitate of double carbonate, easily soluble in excess; from this solution the uranium is reprecipitated by the addition of NaOH or KOH, or by boiling the liquid; it cliffers in this respect from the corresponding iron precipitate.
 - 386. Na₂HPO₄: yellowish-white precipitate of $\rm UO_2HPO_4$; in the presence of AmA, uranyl ammonium phosphate ($\rm UO_2~Am~PO_4$) is precipitated.
 - 387. K_4 FeCy₆ gives, in acid solutions, a reddish-brown precipitate of $(UO_2)_2$ FeCy₆, which differs from the similar precipitate produced in a copper solution by dissolving in excess of AmOH to a *yellow* liquid. This is a very delicate test.
 - 388. BaCO $_3$ causes complete precipitation as uranyl barium carbonate, Ba $_2$ UO $_2$ (CO $_3$) $_3$ even in the cold.
 - 389. Zn changes the yellow colour of acid solutions to green.
 - 390. Borax and Microcosmic beads: after being heated in the outer flame, yellow, and in the inner flame, green.

VANADIUM (V).-Use Sodium vanadate, NaVO3.

- 391. V occurs in vanadinite, $3\text{Pb}_3(\text{VO}_4)_2.\text{PbCl}_2$, and in certain iron-ores and copper-ores; also in many clays and in most granites. Vanadium is known in several stages of oxidation; it is usually found in analysis as vanadic acid or a vanadate, which corresponds to the oxide V_2O_5 . In acid solution the vanadates are yellow or reddish in colour.
- 392. Reducing agents: H_2S , H_2SO_3 , or $H_2C_2O_4$, reduces an acid solution of a vanadate to a divanadyl compound, and causes the colour of the solution to change to *blue*; with H_2S a deposit of sulphur is also formed.
- 393. Zn, when it is introduced into a very dilute solution, which is acidified with $\rm H_2SO_4$ and is then gently warmed, causes the liquid to change in colour to blue, then to green, and then to lavender-blue. The changes of colour are due to successive stages of reduction
- 394. Am_2S , if added in excess, gives a brown liquid containing sulphosalts; the addition of an excess of acid to this brown liquid produces a brown precipitate of V_2S_5 ; the brown precipitate is redissolved by alkaline hydroxides and by Am_2S , forming a reddish-brown solution.
- 395. Solid AmCl, if it is added to the solution until it is saturated, precipitates Am₃VO₄ which is insoluble in saturated AmCl-solution.
- 396. H_2O_2 , when it is shaken up with an acid solution, imparts to it a red colour; the colour is not removed by shaking the liquid with ether. This is a very delicate test.
- 397. Borax-bead: in the outer flame colourless, or yellow if much \hat{V} is present: in the inner flame green hot and cold, or, if much V is present, brown hot and green cold.

DETECTION OF THE RARER ELEMENTS IN GROUP III.

398. For the Detection of the Razer Elements of Group III. in the ordinary course of analysis, refer to the third and fourth columns in the Tables in pars. 1323, 1324.

GROUP IV.—BARIUM GROUP.

399. This Group includes Ba, Sr, Ca.

Analytically these metals differ from those of Group V. by being precipitated as carbonates by Am₂CO₃, and as oxalates by Am₂C₂O₄, in the presence of AmCl. They are not precipitated by any other Group-reagent. The carbonates are soluble in boiling solution of AmCl.

Barium Group of Metals.

400. The metals of this Group are difficult to isolate, and they oxidise rapidly when they are exposed to the air. Their compounds correspond to the oxide MO. Their hydroxides are soluble and alkaline and are strongly basic. The soluble salts which correspond to the strong acids are neutral, and they contain a colourless divalent ion.

The metals are often termed the "metals of the alkaline earths," since the oxides produced from the natural carbonates are alkaline to test-paper when they are dissolved.

REACTIONS OF BARIUM (Ba).

Use Barium chloride, BaCl₂.2H₂O.

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For minerals containing Ba refer to par. 1419.

- 401. Am₂CO₃, added after AmCl, causes a white precipitate (BaCO₃). This precipitate is at first flocculent; but it slowly shrinks in volume and becomes crystalline, if the liquid which contains it is gently heated and is then allowed to stand. Prove that this precipitate is soluble in HĀ.
- 402. CaSO₄, or H₂SO₄, causes a heavy white precipitate (BaSO₄), which is insoluble in HCl even when the liquid is boiled.
- 403. K₂CrO₄, added after some HĀ, produces a yellow precipitate (BaCrO₄), which is soluble in warm HCl, but differs from SrCrO₄ by being insoluble in warm HĀ.
- 404. Hydrofluosilicic acid, H₂SiF₆, produces a semi-transparent precipitate (BaSiF₆). The precipitate does not appear in dilute solutions until after some time; its formation is hastened by boiling, thaking, or stirring the liquid, or by adding alcohol. H₂SiF₆ yields no precipitates with solutions containing Sr or Ca.
 - 405. Ammonium oxalate, Ara C2O4, produces a white precipitate (BaC2O4).

406. Flame-coloration: BaCl₂ gives a yellowish-green colour to the flame, which is visible through the indigo-prism.

The barium-spectrum (Fig. 51, p. 50) consists of a number of lines, the most characteristic of which are three green lines, α , β , γ .

REACTIONS OF STRONTIUM (Sr).

Use Strontium nitrate Sr(NO₃)₂.4H₂O.

For minerals containing Sr refer to par. 1419.

- 407. Am₂CO₃, added after some AmCl, gives a white precipitate (SrCO₃), which is soluble in HĀ. This precipitate is flocculent at first; but it gradually becomes crystalline and shrinks very considerably at ordinary temperatures, and more rapidly when it is heated.
- 408. CaSO₄, or H₂SO₄, gives a white precipitate (SrSO₄). This precipitate does not usually form in a cold solution at once, but only after some time. The precipitate, however, appears at once when the liquid is boiled.
- 409. K_2 CrO₄, added after some HĀ, causes no precipitate, since SrCrO₄ is soluble in HĀ.
 - 410. Am₂C₂O₄ produces a white precipitate (SrC₂O₄).
- 411. Flame-coloration: crimson-red; this colour appears intense red through the indigo-prism, unless the flame-coloration is very faint.

The strontium-spectrum contains many lines (Fig. 51, p. 50). The most characteristic are the orange line (a), the red lines (β, γ) , and the blue line (δ) .

REACTIONS OF CALCIUM (Ca).

Use Calcium chloride, CaCl₂.6H₂O.

For minerals containing Ca refer to par. 1419.

- 412. Am₂CO₃, added after AmCl, gives a white precipitate (CaCO₃) which is soluble in HĀ. This precipitate is flocculent at first; but it shrinks considerably and becomes crystalline after a time, more rapidly when it is gently heated.
- 413. CaSO₄: no precipitate, even when the liquid is allowed to stand, or is boiled.
 - 414. H2SO4: a white precipitate (CaSO4) forms at once in

strong solutions, and often in weak solutions when they are boiled; but some CaSO₄ will always remain dissolved, since it is not quite insoluble in water.

Prove this by boiling the liquid, in which the further addition of H SO₄ has caused no precipitate and which contains the precipitate, and filtering it. Then keep adding AmOH to the filtrate and stirring it, until a drop of the solution turns turnseric-paper brown and the liquid smells of NH₃. Then add Am₂C₂O₄; a white-precipitate will form, showing the presence of Ca (416) in solution as CaSO₄.

415. K2CrO4: no precipitate.

- 416. $Am_2C_2O_4$: white precipitate (CaC_2O_4) , soluble in most acids, but insoluble in $H\bar{A}$ and in oxalic acid, $H_2C_2O_4$.
- 417. Flame-coloration: yellowish-red; when this coloration is viewed through the indigo-prism it does not appear red, and differs in this respect from that given by Sr.

In the calcium-spectrum (Fig. 51, p. 50) the most characteristic lines are the green line (β) and the intense orange line (a).

DETECTION OF THE METALS IN GROUP IV., WHEN THEY OCCUR TOGETHER.

- 418. The following Differences presented by the solutions of these metals are made use of:—
- 1. Their different flame-colorations, which are examined if necessary through the indigo-prism and by means of the spectroscope.
- 2. The difference in the behaviour of their solutions with CaSO₄-solution, which at once indicates the presence or absence of Ba, or of Sr in the absence of Ba.
- 3. The insolubility of BaCrO₄ in HĀ, and hence the possibility of separating Ba, if present, by adding HĀ and then K₂CrO₄.
- 4. The fact that Sr, but not Ca, can be precipitated by CaSO₄ on boiling. This reagent will therefore indicate the presence of Sr in the absence, or after the separation of Ba.
- 5. The complete precipitation of Sr, if present, by boiling the solution with H₂SO₄; sufficient CaSO₄ still remaining in the filtrate to give a precipitate on the addition of AmOH in excess and of Am₂C₂O₄.
- A Solution which has to be Examined for Ba, Sr, and Ca, and which can contain only these metals, is first made alkaline,

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if it is not already so, by the addition of AmOH in excess. Am₂CO₃ is then added, and the addition of this reagent is continued as long as it causes any further precipitate after the liquid has been warmed, stirred well and allowed to settle. The liquid is then filtered.

A little more Am₂CO₃ is now added to the clear filtrate, and if this causes any further precipitate more Am₂CO₃ is added, and the liquid is again poured through the same filter. These processes are repeated, if necessary.

As soon as the filtrate gives no further precipitate with Am₂CO₃, all the metals of this Group, which were present in the solution, will have been precipitated as carbonates, and will remain on the filter after the filtration of the liquid.

The precipitate is then examined by par. 1260, and by Table IV. (1261), or by the simpler Table IV.A. (1262).

GROUP V.—THE POTASSIUM GROUP.

419. This Group includes K, NH₄, Na, Li, Mg, together with the rarer metals Rb, Cs.

The members of this Group are not precipitated from their solutions by any Group-reagent.

The reactions of the rarer metals, Rb and Cs, are given in pars. 446, 447.

The Alkali Metals.

420. Li, Na, K, Rb and Cs are known as the alkali metals. The radicle ammonium is included in this analytical group because its compounds and their reactions closely resemble those of the alkali metals.

• These metals are rapidly oxidised in the air, and are converted ultimately into their peroxides. They decompose water at ordinary temperatures with evolution of hydrogen and formation of strongly basic alkaline hydroxides.

Their compounds correspond to the oxide M₂O, and are usually colourless and soluble. The oxides are strongly basic. The solutions of the salts corresponding to the strong acids are neutral in

reaction, and solutions of salts corresponding to weak acids are alkaline in reaction.

The salts are usually somewhat volatile and impart distinctive colours to a non-luminous flame, which give characteristic and simple spectra.

Note.—The solution of H₂PtCl₆ which is used for the detection of some of these metals is commonly known as platinum chloride (PtCl₄) solution, but it actually contains chlorplatinic acid, H₂PtCl₆; both of these names and formulæ are used in the text.

REACTIONS OF POTASSIUM (K).

Use Potassium chloride, KCl.

For minerals containing K refer to par. 1419.

421. Platinum chloride, or Chlorplatinic acid, H₂PtCl₆ (see *Note* above), if it is added to some of the KCl-solution on a watch-glass, and stirred gently but persistently with a glass rod, gives a yellow crystalline precipitate (K₂PtCl₆); this precipitate forms first along the lines where the rod has rubbed the glass.

Stir up the precipitate and pour off the liquid with the precipitate into three test-tubes.

To one of these portions add much distilled water and heat, the precipitate will dissolve. To another part add alcohol, the precipitate will not dissolve. Hence this precipitate is soluble in water and insoluble in alcohol.

By warming the third portion with solution of potassium hydroxide (KOH), the precipitate is shown to be soluble in that liquid.

Remarks.—Since this precipitate is soluble in alkalis, a liquid, to which this test has to be applied, should always be made neutral or faintly acid before H₂PtCl₆ is added. It is always well to acidify the liquid with dilute HCl, and to make sure by test-paper that it is slightly acid in reaction.

Further, since the precipitate is less soluble in alcohol than in water, the addition of alcohol often hastens the formation of the precipitate in dilute solutions.

Owing to the solubility of the precipitate in water, very dilute solutions must be evaporated nearly to dryness and cooled before the test is applied

422. Sodium picrate, if it is added to a little KCl-solution on a watch-glass, gives golden-yellow needle-like crystals of potassium picrate when the liquid is stirred gently with a glass rod. The potassium solution should be neutral, or faintly alkaline, and free from sammonium salts. The reaction is promoted in a dilute solution by the addition of alcohol; it is not affected by the presence of acetic acid.

423. Sodium hydrogen tartrate, NaHT, if it is added to some of the KCl-solution in a test-tube, gives a white crystalline precipitate (KHT). The precipitate usually appears only when the tube is well shaken for some time, after the mouth of the test-tube has been closed with the thumb.

Shake up the precipitate, and pour it off with the liquid into four test-

Add to three of these different portions, water, KOH, and HCl respectively; on being shaken or warmed with each of these liquids, the precipitate will dissolve.

To the fourth portion add alcohol; the precipitate will not dissolve.

Remarks.—When the above test is used for the detection of potassium, the precipitate will not be obtained at once in dilute solutions; but its formation is much hastened by the addition of alcohol. Very dilute solutions must first be concentrated by evaporation.

The solution to be tested should be neutral or feebly alkaline in reaction. If it is strongly acid in reaction, the free acid should be neutralised by the addition of Na₂CO₃-solution; or, if the acid is volatile, it may be removed by evaporation. Feebly-acid solutions may be at once mixed with NaHT. To an alkaline solution, tartaric acid (H₂T) should be added until the reaction of the liquid is strongly acid.

424. Flame-coloration (62).—Dip a loop of platinum wire, which gives no colour to the flame, into some strong KCl-solution; or moisten the wire, and dip it into some finely-powdered KCl: then hold the loop in the flame.

A pale-lilac or lavender coloration will be seen, if the KCl is pure. Even if the flame is yellow to the naked eye, owing to the presence of Na, it will appear deep *crimson* when it is viewed through the thicker parts of an indigo-prism (Exp. 33, 62).

The flame-coloration due to potassium shows a spectrum of two lines, a red line (a) and a faint indigo-blue line (β); these lines are at the extreme ends of the spectrum (Fig. 51, p. 50): a faint continuous spectrum is also visible.

425. Heat a little Solid KCl in a small dry test-tube; or powder it finely, and heat it on a piece of platinum-foil. The KCl will melt, but it will only produce slight white fumes when it is heated very strongly. This proves that potassium salts are very slightly volatile even at a bright red heat.

The fused residue is often transparent and invisible; but its presence may always be detected by a slight crackling being heard, during the cooling of the foil or of the test-tube.

REACTIONS OF AMMONIUM (NH4).

Use Ammonium chloride, NH4Cl or AmCl.

426. Potassium hydroxide, KOH, if it is poured in excess

into some AmCl-solution, or upon the solid, and heated, causes ammonia gas to be given off:-

$$NH_4CI + KOH = NH_3 + K(1 + H_2O)$$

The ammonia is recognised:-

- (a) By its pungent smell, which is that of common "smelling salts."
- (b) By turning moistened red litmus-paper blue, or turmeric-paper brown.

Wet a small piece of the test-paper with distilled water, and place it along the end of a glass rod. Then hold the rod in the test-tube, taking great care not to bring the paper into contact with the liquid or with the sides of the tube. The change of colour mentioned above will take place.

- (c) By yielding white fumes with an acid gas or vapour. Hence if a glass rod or stopper, wetted with strong H(1, HNO₃, or HĀ, is held over the mouth of the test-tube, dense white fumes will be seen.
- 427. Platinum chloride or Chlorplatinic acid, H₂PtCl₆ (see *Note*, 420), when it is stirred on a watch-glass with some Am(!solution, gives a yellow crystalline precipitate (Am₂Pt(!l₆).

This precipitate is soluble in water and in KOH-solution, but is insoluble in alcohol. If it is boiled with KOH, NH₂ is evolved, and it is distinguished in this way from the similar potassium precipitate, K₂PtCl₆. The remarks in small type after paragraph 421 apply to this reaction also.

- 428. Sodium picrate, when it is stirred on a watch-glass with some AmCl-solution, gives golden-yellow needle-like crystals, which are soluble in much water (see 422).
- 429. Sodium hydrogen tartrate, NaHT, when it is shaken for some time in a test-tube with AmCl-solution, gives a white crystalline precipitate, AmHT.

This precipitate is soluble in water, in KOH, and in HCl, but is insoluble in alcohol. The remarks in small type in paragraph 423 apply to this reaction also.

- 430. Flame-coloration.—Pure AmCl gives no flame-coloration; but since ammonium salts almost invariably contain a trace of sodium, they usually impart a yellow colour to the flame.
- 431. Heat a little Solid AmCl on a piece of platinum-foil: it will be converted into vapour and will produce copious white fumes. These fumes are best seen when the foil is removed for moment from the flame. The solid at last entirely disappears, showing that AmCl is completely volatile.

Repeat the experiment by heating a small piece of solid AmCl in a dry test-tube. The solid may be completely volatilised, and its vapour condensed as a white coating or "sublimate" on the upper part of the tube.

Some ammonium salts, such as the carbonate and nitrate, are decomposed by heat into gases; they are therefore volatilised without producing white fumes or a sublimate.

REACTIONS OF SODIUM (Na).

Use Sodium chloride, NaCl.

For minerals containing Na refer to par. 1419.

432. Flame-coloration.—NaCl gives an *intense yellow* colour to the flame. The colour is either invisible, or appears only pale blue, when it is viewed through the indigo-prism; it never shows the slightest tinge of red when it is seen through the thicker parts of the prism.

When the sodium coloration is examined by a small spectroscope, it gives a single yellow line (α) (Fig. 51, p. 50).

The yellow coloration of the flame which is produced by sodium is usually readily seen, even in the presence of other flame-colorations, since it is not easily masked. In any case the yellow spectrum line may always be detected, even when mere traces of sodium are present.

If both potassium and sodium are present, the potassium coloration is usually unseen, and the yellow coloration of sodium alone is visible. But when the flame is viewed through the indigo-prism, the crimson coloration of potassium is at once seen (62, Exp. 33). The lines of Na and K may also be distinctly separated from one another by means of the spectroscope.

433. Heat a little Solid NaCl in a dry test-tube or on platinumfoil: it will melt without producing white fumes, and will give slight fumes only when it is heated very strongly, showing that NaCl is very slightly volatile even at a bright red heat.

Magnesium (Mg).

Density 1.75, melting-point 750° C., boiling-point 1100° C.

434. Magnesium is a white metal, which slowly oxidises in moist air, and burns brilliantly to its oxide when it is kindled in the air. It is readily soluble in acids and slowly decomposes water with evolution of hydrogen.

The metal resembles the metals of the Barium, or alkaline earths group, (400) in its general chemical characters, but its oxide is less basic. Its compounds correspond to the oxide Mg(), and its soluble salts contain a colourless divalent ion.

For minerals containing Mg refer to par. 1419.

REACTIONS OF MAGNESIUM (Mg).

Use Magnesium sulphate, MgSO₄.7H₂O.

435. Add AmCl, then AmOH, and then Sodium phosphate, Na₂HPO₄: a white crystalline precipitate (MgAmP()₄.6H₂()) will be produced:—

 $\mathrm{MgSO_4} \, + \, \mathrm{AmOH} + \mathrm{Na_2HPO_4} = \mathrm{MgAmPO_4} + \mathrm{Na_2SO_4} + \mathrm{H_2O}.$

If much water is present, this precipitate will form only when the liquid is warmed, or is well stirred or shaken: it is soluble in HCl and in other acids, but is insoluble in AmOH.

- 436. KOH gives a white precipitate, $Mg(OH)_2$, which is readily soluble in acids.
- 437. AmOH gives a white precipitate, $Mg(OH)_2$; but if AmCl is added before the AmOH, no precipitate is produced.
- 438. $\rm Am_2CO_3$ gives a white precipitate, MgCO₃, but only in strong solutions and on standing for some time. The previous addition of AmCl prevents the formation of this precipitate.
- 439. Flame-coloration.—Pure MgSO₄-solution gives no colour to the flame.
- 440. Blowpipe-test.—If a little solid magnesium sulphate is placed in a small cavity on a piece of wood-charcoal, and is then heated in the tip of the outer blowpipe-flame, it will shine brightly and will give a white unmelted mass of MgO. No white fumes will be seen, since MgO is not volatile.

If this white mass is allowed to cool and is moistened with several drops of cobalt nitrate solution, and is then again heated to whiteness for some time in the outer blowpipe-flame, it will assume a delicate pink colour. This colour will be readily seen when a piece of white paper is held near the cold residue on the charcoal.

Reactions of Lithium (Li). .

Use Lithium chloride, LiCl.

At Lt occurs frequently in mineral waters and in the ashes of plants, also in small quantities in the minerals lepidolite, triphane, and petallite: the minerals lepidolite, triphane, and petallite:

Li salts resemble salts of the metals of the Barium Group by the difficult solubility of the carbonate and phosphate; they differ from those of K and NH₄ by not being precipitated by H₂PtCl₆ or by NaHT; Li is readily distinguished from Na by its flame-coloration and its spectrum.

LiCl may be separated from KCl and NaCl by its insolubility in a mixture

of equal volumes of absolute alcohol and dry ether.

442. Sodium phosphate, Na₂HPO₄, if it is added to the solution, which must not be too dilute and must be made strongly alkaline with NaOH, gives on boiling a white crystalline precipitate (Li₃PO₄) which settles quickly.

Traces of Li may be precipitated by adding Na₂HPO₄, and then NaOH until the liquid remains alkaline, evaporating to dryness and washing the

residue with dilute AmOH.

This precipitate may be distinguished from the phosphates of Ba, Sr, Ca, and Mg by heating it on charcoal in the blowpipe-flame, when it will readily

fuse and be absorbed by the charcoal support.

The diluted cold solution of Li₃PO₄ in HCl also gives no precipitate when AmOH is added in excess, but a white crystalline precipitate forms when the liquid is boiled. Also the phosphate precipitate, when it is moistened with HCl, gives the characteristic flame-coloration and spectrum (443).

443. Flame-coloration: this is carmine-red; the spectrum (Fig. 51, p. 50) consists of an intense carmine-red line (a)

The flame-coloration is concealed by that of Na, but the presence of Na does not interfere with the spectrum, and the Na-coloration may be removed by viewing the flame through the thinner parts of the indigo-prism.

The lithium flame-coloration differs from that given by K by being either unable to penetrate the thick layers of indigo-solution in the indigo-prism, or by being much lessened in intensity by passage through the indigo.

Lithium silicate only gives the flame-coloration after it has been fused with CaSO₄; lithium phosphate requires to be first moistened with HCl.

DETECTION OF METALS IN GROUP V.

- 444. The detection of metals of this Group, when they occur together, depends upon the following differences:—
- 1. Mg is indicated with certainty by its precipitation, even in dilute solutions, by $\rm Na_2HPO_4$ in the presence of AmCl and AmOH, and by its precipitation by KOH-solution.
- 2. Na can be found by its yellow flame coloration; and K may be found by its lilac flame coloration, or in the presence of Na by a crimson flame-coloration being seen through the indigo-prism, or by its spectrum.
- 3. Li may be detected by its carmine-red flame coloration, which is not readily seen through the indigo-prism, or in the presence of Na by its spectrum.
- 4. NH₄ may be detected by the evolution of NH₃ when the substance is boiled with KOH-solution.
- 5. The test for K'by means of H₂PtCl₆, NaHT, or sodium picrate is only conclusive in the absence of NH₄. If NH₄ is present, therefore, it must be removed by ignition before K can be tested for by precipitation.

It will be found more convenient to use the scheme in paragraph 445 than the Table given in paragraph 1263, and it will generally

be unnecessary to try the confirmatory test (IV.) for K, since K may be detected with certainty by its flame-coloration and by its spectrum.

Sodium is rarely absent from any substance. The quantity of this metal which is present may be judged from the flame-coloration: unless the colour is intense, only a trace of sodium is present.

Lithium has been included in this scheme because its compounds are of fairly common occurrence, but it is usually classed amongst the rarer metals of the Group.

SCHEME FOR DETECTING K, NA, LI, NH₄, MG WHEN THEY MAY OCCUR TOGETHER IN SOLUTION.

- 445. The scheme includes the following three tests or experiments, and a fourth confirmatory test may also be made. If a solid is to be examined it may be dissolved in water, or if necessary in dilute HCl.
- I. Dip a clean loop of platinum wire into the solution and hold it near the top of the Bunsen-flame, watching for some minutes to see if any initial coloration undergoes change. Note the flame coloration, and if necessary confirm its indications by observation of the spectrum (Fig. 51, p. 50):—

A lilac coloration indicates Presence of K.

A crimson coloration shows Presence of Li.

A yellow coloration proves Presence of Na, and if the colour appears deep crimson through the indigo-prism this shows also Presence of K.

II. Add to a small part of the solution, after dilution if Li has been found (442), AmCl, AmOH and Na₂HPO₄, stir the liquid well and allow it to stand

A white precipitate forms, showing Presence of Mg.

III. Boil some of the solution with excess of K()H-solution, and try whether NH_s is evolved, by smelling, or if necessary by moist litmus-paper or strong HCl introduced on a glass rod (426): NH_s is smelt, or the turmeric becomes brown, or white fumes are produced with the HCl, indicating Presence of NH_s.

[If Mg is absent a confirmatory test for Li may be made by adding Na, HPO4 to the above boiling alkaline liquid, when white Li, PO4

will be precipitated.]

IV. The presence of K may be confirmed, if NH₄ is absent, by precipitating another part of the solution with H₂PtCl₈ (421), sodium picrate (422) or NaHT (423).

If NH₄ is present it must be removed by evaporating some of the solution to dryness in a porcelain dish, igniting the residue on platinum-foil until no more fumes are evolved when the foil is removed from the flame, dissolving the residue by boiling the foil in a test-tube with a little water, and testing this solution, after concentration if necessary by evaporation, with one of the above reagents for detecting K (421-423).

REACTIONS OF THE RARER METALS IN GROUP V.

The rarer metals included in this Group are Rb, Cs.

RUBIDIUM (Rb), CÆSIUM (Cs).—Use RbCl and CsCl.

446. The metals Rb, Cs occur in small quantities in some mineral waters, in carnallite, and in a few minerals.

The compounds of Rb and Cs resemble those of K by being precipitated by $\rm H_2PtCl_6$ and by $\rm H_2\overline{T}$, and by giving a flame-coloration similar to that of K.

These metals differ from K by the greater insolubility in water of their chlorplatinates. When these have been precipitated together, K₂PtCl₆ can be dissolved away from Rb₂PtCl₆ and Cs₂PtCl₆ by boiling the precipitate with successive small quantities of water. The alums containing these metals show a similar difference in solubility in cold water.

447. Flame-colorations.—The flame-colorations given by these metals are not distinguishable from that given by K. Cs and Rb are, however, readily distinguished from one another and from other elements by their spectra (Fig. 51, p. 59).

The blue lines (α, β) , given by Cs, are especially distinct and characteristic. In the Rb-spectrum the indigo-blue lines (α, β) are very distinct, but the red lines (γ, δ) are most characteristic.

The chlorides are the most suitable salts for the spectroscope test.

448. The metals Li, Rb, Cs are most readily detected in this Group by means of their spectra.

For the detection of these metals in the ordinary course of analysis refer to the fifth column of the Table in paragraph 1323.

SECTION IV.

REACTIONS OF THE INORGANIC ACID-RADICLES.

460. Remarks on Acids and Acid-radicles.—Acids may be defined as compounds, which are resolved by solution in water into the positive ion H, and a negative ion called the "acid-radicle" (81): the acid-radicle may be either an element or a group of elements. Thus HCl and HNO₃ are resolved by solution in water into H' + Cl' and H' + (NO₃)' ions respectively.

The replacement of the hydrogen in an acid by a metal produces a "salt."

A "strong" acid is ionised to a large extent when it is dissolved; a "weak" acid is ionised only to a small extent by solution.

Sulphuric acid, nitric acid, the halogen acids (except hydrofluoric) and a few others are the common strong acids. They yield a large number of metallic salts, and the salts which are formed by the action of these strong acids upon the strong bases are neutral in reaction.

The weak acids, such as hydrogen sulphide, carbonic acid, silicic acid, and boric acid yield fewer salts, and the salts formed by their action upon the strong bases are alkaline in reaction.

There are also acids of intermediate strength, such as phosphoric acid and sulphurous acid; the salts corresponding to these acids are also of an intermediate character.

- 461. Trying Reactions.—The reactions for the Acid-radicles are tried in the same way as those for the metals (83-87), a salt or other compound containing the radicle being employed. A salt of sodium, potassium or ammonium is usually preferred for this purpose, since these metals do not as a rule interfere with the reactions of their associated Acid-radicles.
- 462. Acid-radicle Groups.—The Acid-radicles are usually detected in the course of analysis by special tests. They cannot advantageously be precipitated in Groups, from which the members of each Group are subsequently separated and identified, as is the ease with the Metals.

Accordingly the classification adopted for their reactions consists in placing together in a Group those Acid-radicles which some-

what resemble one another in their reactions; and then stating at the end of each Group the differences upon which the detection of its members depends, when the radicles occur together.

At the end of each Group the student may with advantage detect one or more of its members in an unknown substance, as he has already done with the metals.

GROUP I.—SULPHATE GROUP.

463. This Group includes sulphate (SO₄), selenate (SeO₄), and fluosilicate (SiF₆).

These acid-radicles are distinguished by yielding a precipitate, when ${\rm BaCl}_2$ is added in the presence of hydrochloric acid.

Sulphate is the only commonly occurring acid-radicle which gives with BaCl₂ a precipitate insoluble in boiling HCl.

Fluosilicate resembles sulphate by its Ba-precipitate being only slightly soluble in boiling HCl, but it is also related to silicate and fluoride (645-657).

Selenate yields a precipitate of BaSeO₄ when BaCl₂ is added in the presence of cold HCl, but this precipitate differs from BaSO₄ and BaSiF₆ by being dissolved when it is boiled for some time with strong HCl (252).

SULPHATE, (SO₄)".—Use Na₂SO₄.10H₂O.

464. BaCl₂, or Ba(NO₃)₂; white precipitate, BaSO₄, which remains undissolved when it is boiled with dilute HCl or HNO₃.

Note.—BaCl₂ and Ba(NO₃)₂ are themselves insoluble in strong HCl and in strong HNO₃. Care must therefore be taken not to mistake precipitates caused by these acids for a precipitate of BaSO₄. These precipitates are readily distinguished from BaSO₄ by their solubility in water.

Add a little BaCl₂- or Ba(NO₃)₂-solution to strong HCl or HNO₃. A white precipitate will be produced, consisting of BaCl₂ or Ba(NO₃)₂, which is insoluble in the strong acid. Now add much water, the precipitate will dissolve readily and completely in the water.

465. Blowpipe-test on Charcoal.—If a sulphate, or any solid substance containing sulphur, is mixed with solid Na₂CO₃, and the mixture is then heated on charcoal in the inner blowpipe-flame until the Na₂CO₃ fuses, Na₂S will be produced:—

$$Na_2SO_4 + 4C = Na_2S + 4CO$$
.

The Na₂S thus produced may be readily detected by detaching the cold solid mass from the charcoal with the point of a knife, placing a portion of it upon a bright silver coin, and moistening it with a drop of water: after the moistened mass has remained upon the coin for a short time, it is rinsed off, a black stain (Ag₂S) will then be seen upon the part of the silver upon which the substance rested (481).

Another portion of the cool mass may be moistened with a drop of HCl: a fetid smell of H_2S will be observed, and a piece of paper moistened with $Pb\bar{A}_2$ -solution and held above the substance will be blackened (480).

This test is evidently only of value for detecting a sulphate, when it is known that sulphur is not present in any other form in the substance to be tested.

It is also necessary that sulphur should not be introduced by the Na₂CO₃, or by the flame. Since coal-gas may contain sufficient sulphur to give the above reaction with pure Na₂CO₃, it is advisable to employ the flame of a spirit- or petroleum-lamp (5, 6), rather than that of a gas-burner, for this reaction.

466. Hydrogen sulphate, or Sulphuric acid, H₂SO₄, in the cold undiluted condition, slowly chars many organic substances: this change is rapid when the acid is heated. If the acid is dilute it may be detected by producing black charred stains on filterpaper, which has been spotted or marked with the liquid and has then been dried by heat.

Sulphuric acid has a strongly acid reaction to litmus, and evolves hydrogen when it is diluted and warmed with Zn.

The above properties are, however, also shown by some metallic sulphates and by the acid sulphates.

SELENATE (SeO₄)".—The reactions for selenate have been already given (251-254) in Group II. for the metals. Selenate is classed in Group II. because it gives a precipitate with H₂S, after its solution in hydrochloric acid has been boiled.

SILICOFLUORIDE, or FLUOSILICATE, $(SiF_6)''$.—Use Hydrogen silicofluoride or Hydrofluosilicie acid, H_2SiF_6 .

467. BaCl₂: white precipitate, BaSiF₆: this precipitate usually appears only when the solution is gently warmed, and differs in this respect from BaSO₄. It differs from BaSeO₄ (252) by being only slightly soluble in boiling HCl: it is insoluble in alcohol.

Allow this precipitate to settle and reserve it for the reaction in paragraph

The above precipitate shows some resemblance to BaSC₄ since it is only slightly soluble in HCl, but it differs in appearance by being more or less transparent and crystalline. Silicofluorides differ also from sulphetes by

giving no precipitate with $Sr(NO_3)_2$ or with $Pb(NO_3)_2$. Further, when their solutions are heated with excess of AmOH, they yield a flocculent precipitate of H_2SiO_3 and leave AmF in solution (655).

Silicofluorides also differ from sulphates by the following very distinctive

reactions.

468-472]

468. KCl-solution, when it is warmed with solution of silicofluoride, gives a very transparent gelatinous precipitate, $K_2 SiF_6$: this precipitate is only visible after it has been allowed to settle for a time: it is insoluble in alcohol.

469. Strong \dot{H}_2SO_4 , if it is heated with a silicofluoride, causes HF to be given off, which may be detected by its etching action on glass (654).

This reaction is especially useful for ascertaining whether a precipitate, which has been produced by BaCl₂, consists of or contains BaSiF₆, and the precipitate from paragraph 467 may be tested by this process, as follows:—

The precipitate is allowed to settle: almost all the water is then poured off: the remainder of the water is shaken up with the precipitate and is at once poured into a watch-glass: the remaining water is then decanted, and the precipitate which remains on the glass is warmed for some time with strong $\rm H_2SO_4$. After the glass has been washed and dried, it will be found to be corroded where the precipitate has rested.

The more delicate process described in par. 654 may be substituted, the precipitate, separated by decantation or by filtration and drying, being

treated in a platinum or leaden cup.

470. CaCl₂ gives no precipitate. [Difference from fluoride (656).]

471. Hydrogen silicofluoride, or Hydrofluosilicic acid, H_2SiF_6 , is a strongly acid liquid, which volatilises completely as $SiF_4 + 2HF$ when it is heated in platinum, and therefore etches a watch-glass if it is evaporated to dryness in the glass. It gives reactions 467 and 468.

DETECTION OF ACID-RADICLES IN GROUP I.

472. After the acid-radicle has been precipitated by the addition of BaCl₂, it may be identified by noting the following differences in the properties of the barium precipitate.

The precipitate of BaSO₄ differs from that of BaSeO₄ by being

quite unaffected when it is boiled with HCl.

BaSeO₄ may be dissolved away from BaSO₄ by being boiled with HCl, and selenite may then be detected in solution by the tests given in paragraphs 246-250. When BaSeO₄ is heated on charcoal, it gives the peculiar smell of rotten horse-radish (253).

BaSiF₆ is distinguished by heating it with strong H₂SO₄, when HF is evolved, and etches glass (469). For other differences, see

paragraphs 467, 468.

ž .

GROUP II.—CARBONATE GROUP.

473. This Group includes carbonate (CO_3) , sulphide (S), sulphite (SO_3) , thiosulphate (S_2O_3) , hypochlorite (CIO), and nitrite (NO_2) .

Salts which contain these acid-radicles evolve characteristic gases when they are acted upon by dilute HCl.

CARBONATE, (CO₃)".—Use Na₂CO₃, or CoO₃.

474. HCl, $H\bar{A}$, $H_2\bar{T}$, and almost any other acid, when it is poured upon a carbonate in a test-tube, causes colourless, almost inodorous carbon dioxide gas, CO_2 , to escape with effervescence. This gas is recognised by its property of turning lime-water or baryta-water milky, owing to the production of an insoluble carbonate.

The Lime-water test may be tried in four ways, as is described below, attention being paid to the precautions which follow.

- 1. A glass rod which has been just removed from some perfectly clear lime-water, and which has a small drop adhering to its end, is introduced into the test-tube in which the carbon dioxide has been evolved; the drop will quickly become milky. Care must be taken not to allow the rod to touch the liquid or the inside of the tube. The milkiness is best seen when the rod is removed and the drop is held above a dead black surface.
- 2. Since carbon dioxide gas is much heavier than air, it will remain in the test-tube in which it has been liberated, if the tube is held crect and its mouth is loosely closed with the thumb. If the tube is then gradually inclined, the heavy gas may be poured into another test-tube containing lime-water, without allowing any liquid to flow out. On closing this tube with the thumb, and shaking the lime-water up with the gas, the liquid will become milky.
- 3. The acid may be poured upon the carbonate in a small beaker, which is immediately covered with a watch-glass the under surface of which has a drop of clear lime-water adhering to the middle: the drop will become milky.
- 4. If the CO₂ is given off in considerable quantity, it may be conducted into some lime-water contained in a separate tube. This is effected by either of the forms of apparatus which are shown in Fig. 56.

Precautions.—In testing for carbonate the following precautions must be attended to.

The milkiness, which is at first produced in the lime-water, gradually disappears when more of the gas is absorbed. Hence, if the lime-water does not become milky at once, it must be constantly watched, in order to avoid the risk of the milkiness not being observed before it has been removed by the further absorption of the gas.

If the addition of the first few drops of acid to the substance does not cause an effervescence, more acid should be added, since a carbonate may

be simply converted into an acid carbonate by the action of a small quantity of acid, and no gas is evolved during that change.

If much water is present in relation to the carbonate, effervescence may not occur, since the gas may remain in solution: it may usually be removed from solution and detected by warming the liquid.



Fig. 56.

APPARATUS FOR TESTING GAS EVOLVED.

A powdered solid substance, which GAS EVOLVED.

is being tested for a carbonate by HCl, should be first moistened with a few drops of water in order to drive out the air-bubbles entangled within it, since if these exaped on the addition of the acid they might be mistaken for a slight effervescence.

475. BaCl₂, or CaCl₂ produces in a solution of carbonate a white precipitate, BaCO₃ or CaCO₃, which is soluble in dilute HCl or HNO₃ with effervescence (474).

ACID OR HYDROGEN CARBONATE, OR BICARBONATE.

476. A Bicarbonate is changed by heat into a normal carbonate with evolution of CO_2 and $H_2O:-2NaHCO_3=Na_2CO_3+CO_2+H_2O$.

Prove this by heating some sodium bicarbonate (NaHCO₃) in an ignitiontube; drops of water will condense, and CO₂ may be detected by lime-water (474).

Also drop some NaHCO₃ into boiling water in a test-tube; effervescence will take place, and the escaping gas will give rise to milkiness in lime-water.

- 477. MgSO₄.—A solution of bicarbonate, which has been made with cold water, gives no precipitate with MgSO₄-solution; but after the solution has been boiled it will contain a normal carbonate, and this will give a precipitate of white MgCO₃.
- '478. HgCl₂-solution yields with solution of bicarbonate a red precipitate; solution of normal carbonate yields a yellow precipitate with HgCl₂-solution.
- 479. Hydrogen carbonate, or Carbonic acid $(\mathrm{H_2CO_3})$, can only exist in dilute aqueous solution, and the addition of sufficient lime-water to its solution causes a permanent milkiness.

Other substances in solution, however, yield a similar result; hence it is best to boil the liquid, and to pass the escaping mixture of steam and CO_2 into lime-water; the lime-water will become milky. Bicarbonates give the same result (476), but a residue of carbonate remains in the solution and may be detected by par. 474, or may be obtained by evaporation.

SULPHIDE, (S)".—Use solid FeS, or solution of Am2S.

480. HCl, or H₂SO₄, when it is poured upon many sulphides, produces H₂S, which may escape with effervescence or may partly

GROUP II.—CARBONATE GROUP.

473. This Group includes carbonate (CO₃), sulphide (S), sulphite (SO₃), thiosulphate (S₂O₃), hypochlorite (ClO), and nitrite NO₂).

Salts which contain these acid-radicles evolve characteristic

gases when they are acted upon by dilute HCl.

CARBONATE, (CO₃)".—Use Na₂CO₃, or CoO₃.

474. HGl, HĀ, H₂T̄, and almost any other acid, when it is soured upon a carbonate in a test-tube, causes colourless, almost nodorous carbon dioxide gas, CO₂, to escape with effervescence. This gas is recognised by its property of turning lime-water or paryta-water milky, owing to the production of an insoluble carbonate.

The Lime-water test may be tried in four ways, as is described below, attention being paid to the precautions which follow.

- 1. A glass rod which has been just removed from some perfectly clear ime-water, and which has a small drop adhering to its end, is introduced nto the test-tube in which the carbon dioxide has been evolved; the drop vill quickly become milky. Care must be taken not to allow the rod to ouch the liquid or the inside of the tube. The milkiness is best seen when he rod is removed and the drop is held above a dead black surface.
- 2. Since carbon dioxide gas is much heavier than air, it will remain in the est-tube in which it has been liberated, if the tube is held erect and its mouth s loosely closed with the thumb. If the tube is then gradually inclined, he heavy gas may be poured into another test-tube containing lime-water, vithout allowing any liquid to flow out. On closing this tube with the humb, and shaking the lime-water up with the gas, the liquid will become milky.
- 3. The acid may be poured upon the carbonate in a small beaker, which is immediately covered with a watch-glass the under surface of which has a drop of clear lime-water adhering to the middle: the drop will become milky.
- 4. If the CO_2 is given off in considerable quantity, it may be conducted nto some lime-water contained in a separate tube. This is effected by either of the forms of apparatus which are shown in Fig. 56.

Precautions.—In testing for carbonate the following precautions must be attended to.

The milkiness, which is at first produced in the lime-water, gradually disappears when more of the gas is absorbed. Hence, if the lime-water does not become milky at once, it must be constantly watched, in order to avoid the risk of the milkiness net being observed before it has been removed by the further absorption of the gas.

If the addition of the first few drops of acid to the substance does not cause an effervescence, more acid should be added, since a carbonate may

be simply converted into an acid carbonate by the action of a small quantity of acid, and no gas is evolved during that change.

If much water is present in relation to the carbonate, effervescence may not occur, since the gas may remain in solution: it may usually be removed from solution and detected by warming the liquid.

A powdered solid substance, which



Apparatus for Testing Gas Evolved.

A powdered solid substance, which is being tested for a carbonate by HCl, should be first moistened with a few drops of water in order to drive out the air-bubbles entangled within it, since if these excaped on the addition of the acid they might be mistaken for a slight effervescence.

475. $BaCl_2$, or $CaCl_2$ produces in a solution of carbonate a white precipitate, $BaCO_3$ or $CaCO_3$, which is soluble in dilute HCl or HNO_3 with effervescence (474).

ACID OR HYDROGEN CARBONATE, OR BICARBONATE.

476. A Bicarbonate is changed by heat into a normal carbonate with evolution of CO_2 and H_2O :— $2NaHCO_3 = Na_2CO_3 + CO_2 + H_2O$.

Prove this by heating some sodium bicarbonate (NaHCO₃) in an ignition-tube; drops of water will condense, and CO₂ may be detected by lime-water (474).

Also drop some NaHCO₃ into boiling water in a test-tube; effervescence will take place, and the escaping gas will give rise to milkiness in lime-water.

477. MgSO₄.—A solution of bicarbonate, which has been made with cold water, gives no precipitate with MgSO₄-solution; but after the solution has been boiled it will contain a normal carbonate, and this will give a precipitate of white MgCO₃.

478. HgCl₂-solution yields with solution of bicarbonate a red precipitate; solution of normal carbonate yields a yellow precipitate with HgCl₂-solution.

479. Hydrogen carbonate, or Carbonic acid (H_2CO_3) , can only exist in dilute aqueous solution, and the addition of sufficient lime-water to its solution causes a permanent milkiness.

Other substances in solution, however, yield a similar result; hence it is best to boil the liquid, and to pass the escaping mixture of steam and CQ_2 into lime-water; the lime-water will become milky. Bicarbonates give the same result (476), but a residue of carbonate remains in the solution and may be detected by par. 474, or may be obtained by evaporation.

SULPHIDE, (S)".—Use solid FeS, or solution of Am S.

480. HCl, or H₂SO₄, when it is poured upon many sulphides, produces H₂S, which may escape with effervescence or may partly

or entirely remain in solution until the liquid is heated. This gas is detected by its fetid smell, resembling that of rotten eggs, and by its property of darkening PbĀ₂-solution by formation of PbS. or of turning nitroprusside solution purple (492).

The gas may be made to act upon the $Pb\bar{A}_2$ -solution by the methods given for CO_2 (474, 1–4); or a small strip of filter-paper, moistened with the solution, may be introduced into the gas on the end of a glass rod. The paper strip should be attached to the rod by only half its length, so as to expose both surfaces of the lower part of the strip to the action of the gas.

Note.—This test is made much more delicate, if the filter-paper is moistened with a solution prepared by adding KOH to boiling $Pb\bar{A}_2$ -solution, until the precipitate which is formed at first is just redissolved. The test-paper which has been thus prepared, becomes intensely blackened by H_2S .

481. Ag.—If solution of sulphide is dropped upon a bright silver coin, it produces a black stain, Ag₂S. This stain may be removed by rubbing the coin with moist lime.

A solid sulphide, which gives off H₂S on the addition of HCl, produces a black stain when it is placed upon a silver coin and is moistened with a drop of HCl.

- 482. PbĀ₂-solution gives with solution of sulphide a black precipitate, PbS. The use of the solution produced by adding KOH in excess to PbĀ₂-solution (480, *Note*) makes this test more delicate.
- 483. Sodium nitroprusside, Na₂FeCy₅NO₂, gives a purple colour even with a mere trace of dissolved sulphide; no colour is given with free H₂S, hence the solution should be made alkaline with AmOH or an alkaline hydroxide before the reagent is added.
- 484. The formation of Thiocyanate (694) may be employed as a very delicate test for a soluble sulphide. It depends on the detection by means of FeCl₃ of the thiocyanate, which has been formed by the action of KCN-solution upon the sulphide.
- 485. Ignition.—Free sulphur and many sulphides, when they are strongly heated in a test-tube, give a sublimate of sulphur in the form of brown drops or of yellow powder.
- 486. Ignition in the Air.—Free sulphur and many sulphider, when they are heated in a tube open at both ends and held obliquely in the flame, produce SO_2 gas. This gas may be recognised by its pungent smell, and by causing the colour of paper moistened with $K_2Cr_2O_7$ -solution to change to bluish-green (490).
- 487. Insoluble Sulphide, when fused with Na₂CO₃ in a covered porcelain crucible or in a hard glass ignition-tube, yields

soluble Na₂S, which may be detected by the above tests (480-483).

- 488. Fusion with KNO₃, or simply heating with strong HNO₃, converts a sulphide into a sulphate, which may be detected in solution by BaCl₂ (464).
- 489. Hydrogen sulphide, or Hydrosulphuric acid (H₂S), either as gas or in aqueous solution, is easily recognised by its smell and by its action on lead acetate solution (480), or it may be mixed with AmOH-solution in excess and the liquid tested by pars. 481-484.

490. HCl, or H_2SO_4 , when it is poured upon a sulphite and warmed, evolves SO_2 gas: this gas may be recognised by its pungent smell and by turning $K_2Cr_2O_7$ -solution green.

A drop of the $K_2Cr_2O_7$ -solution, hanging on the end of a glass rod, may be exposed to the gas (474, 1); or a small strip of filter-paper may be wetted with the bichromate solution and then attached by half its length to the rod, the other half hanging freely in the gas.

491. BaCl₂: white precipitate, BaSO₃: the formation of this precipitate is prevented by the presence of HCl, but a precipitate usually forms because the sulphite often contains traces of a sulphate.

When Cl-water or Br-water is added to the clear solution of ${\rm BaSO_3}$ in HCl, ${\rm BaSO_4}$ is formed and precipitated:—

$$BaCl_2 + H_2SO_3 + H_2O + Cl_2 = BaSO_4 + 4HCl.$$

- . 492. Nascent Hydrogen.—The addition of sulphite solution to a mixture of HCl and Zn, which is giving off pure H, immediately causes an evolution of H₂S. The H₂S is detected by its smell, or by blackening a piece of filter-paper moistened with PbĀ₂-solution, or with Pb(OH)₂ dissolved in excess of KOH (480, Note), or with alkaline nitroprusside solution (492).
- 493. Hydrogen sulphite, or Sulphurous acid, H₂SO₃, exists only in solution. It may be detected by adding to the liquid a drop of K₂Cr₂O₇-solution, when the reddish colour of the dichromate will at once change to green.

Strong aqueous solution of hydrogen sulphite slowly evolves SO₂, and dilute solutions evolve SO₂ when they are heated: the escaping gas may be readily detected by its smell and by K₂Cr₂O₇-solution (490).

Since $\rm H_2SO_3$ -solution is readily oxidised to $\rm H_2SO_4$ by contact with dissolved oxygen, its solution usually contains $\rm H_2SO_4$ unless it has been recently prepared.

Thiosulphate, $(S_2O_3)''$.—Use solution of $Na_2S_2O_3.5H_2O$.

'494. HCl, or H_2SO_4 , causes no immediate change in the cold solution; but after a few seconds the acidified liquid becomes turbid, owing to the separation of finely-divided yellow S, and it then contains $H_2SO_3 := H_2S_2O_3 = S + H_2SO_3$.

SO₂ from the H₂SO₃ may be recognised by its pungent smell, when the acidified liquid is heated; and it may be detected in the solution, by adding a few drops of K₂Cr₂O₇-solution, which changes colour to green.

495. FeCl₃ gives a reddish-violet colour with solution of thiosulphate: this colour disappears after a short time in the cold, and immediately when the liquid is heated, the FeCl₃-solution at the same time loses its yellow colour since it is changed into FeCl₂.

496. AgNO₃ gives a white precipitate, Ag₂S₂O₃, which becomes black after standing for a short time, or immediately when the liquid is heated, owing to the formation of Ag₂S:—

$$Ag_2S_2O_3 + H_2O = Ag_2S + H_2SO_4.$$

The above precipitate, $Ag_2S_2O_3$, dissolves very readily in excess of the $Na_2S_2O_3$ -solution, hence it is most easily obtained by dropping the thiosulphate solution into some $AgNO_3$ -solution.

497. Hydrogen thiosulphate, or Thiosulphuric acid $(H_2S_2O_3)$, is extremely unstable, rapidly separating into $S+H_2SO_3$ (494).

Hypochlorite, (ClO)'.—Use solution of Bleaching-powder, containing Ca(ClO)₂.

498. HClO is evolved from a hypochlorite when it is in the solid state or in strong solution; it is probably liberated by the action of the atmospheric carbon dioxide. The smell of the vapour thus evolved resembles that of Cl, but is less irritating to the nose.

499. HCl, or H₂SO₄, if it is added in excess, sets free Cl in the cold:—

$$Ca(ClO)_2 + 4HCl = 2Cl_2 + CaCl_2 + 2H_2O.$$

The chlorine gas may be recognised by its smell, by its yellowish solener, and by its power of bleaching a piece of moistened litmuster when the paper is held for a short time in the case (474)

- 1-4). If the Cl is small in quantity and remains dissolved, a piece of litmus-paper may be shaken up with the solution.
- The bleaching of litmus is gradually produced by solution of hypochlorite, even without the addition of acid, by the nascent oxygen which is slowly liberated from the hypochlorite. When the hypochlorite solution is heated, or the litmus-paper moistened with the solution is exposed to the carbon dioxide of the air, the bleaching is hastened. On the addition of an acid, however, the bleaching of the litmus-paper immersed in the solution is extremely rapid.
- 500. $Pb\bar{A}_2$: if solution of hypochlorite is added in large quantity to $Pb\bar{A}_2$ -solution, it gives a white precipitate; this precipitate becomes red and then *dark brown* (PbO_2) when the liquid is boiled for a short time.

When hypochlorite is heated with MnCl₂-solution, it also gives a dark brown precipitate of hydrated MnO₂.

501. Hydrogen hypochlorite, or Hypochlorous acid, HClO, is a yellow liquid with sweetish smell; it is very unstable, unless it is largely diluted. This acid bleaches litmus, and evolves Cl when it is warmed with HCl.

Nitrite, (NO₂)'.—Use KNO₂.

- 502. Dilute H_2SO_4 , when it is warmed with nitrite, causes a reddish-brown gas with a characteristic smell to be evolved. The coloured gas is usually best seen by looking down the test-tube at a white surface.
- 503. Cold FeSO₄-solution, when it is added to solution of nitrite, causes the liquid to assume a dark colour: this colour becomes much more intense when cold dilute H₂SO₄ is added.

The dark coloured substance which is produced is a compound of NO with FeSO₄; the compound is decomposed when the liquid is boiled, and reddish-brown gas is evolved (502).

504. Nitrous acid liberates Iodine from Potassium iodide. The nitrous acid may be formed by the addition of acid to the nitrite.

Drop a little starch-powder into some cold water in a porcelain dish, and stir and boil the liquid for a short time: then cool the starch-solution, and add to it a few drops of KI-solution.

Add to this liquid some solution of nitrite, and then several drops of $H\bar{A}$: an intense blue colour will be produced. This is due to the combination of the starch with the iodine, which has been set free by the HNO_3 .

The liquid will often appear black at first, and the blue colour will then only become visible when much water has been added.

This is a most delicate test if the liquid is quite cold. But the

presence of a nitrite is not proved if the colour is produced before the $H\bar{A}$ is added.

505. Metaphenylene diamine gives a very intense yellow coloration when it is added to a slightly acidified solution of a nitrite. This reaction is delicate and is not given by a nitrate.

506. Hydrogen nitrite, or Nitrous acid (HNO_2) , is extremely unstable and decomposes rapidly into HNO_3 , NO, and H_2O . Its solution therefore evolves reddish-brown gas when it is heated in the air (502). Nitrous acid also colours $FeSO_4$ -solution (503), and liberates iodine from KI (504).

DETECTION OF THE ACID-RADICLES IN GROUP II.

507. The Detection of these Acid-radicles, when they occur

singly, presents no difficulty.

Their detection, when they occur together, is often complicated, since most of the acids, which are formed on the addition of HCl to salts containing acid-radicles of this Group, react upon one another. Thus HClO oxidises HNO₂, H₂S, and H₂SO₃; and H₂SO₃ oxidises H₂S.

The analyst may often succeed, by a skilful use of the above reactions, in detecting these acid-radicles when they are mixed. But their detection in certain mixtures is a problem which is only soluble by careful consideration, and is not unfrequently insoluble.

 SO_2 may be detected by $K_2Cr_2O_7$ -paper: H_2S by lead acetate-paper: Cl by bleaching moist litmus-paper: nitrite by evolving reddish-brown gas. But CO_2 can usually only be detected in the presence of much SO_2 after removing the SO_2 : this may be effected by passing the gases through hot $K_2Cr_2O_7$ -solution, when the SO_2 will be absorbed, and the CO_2 which passes on may then be detected by lime-water.

GROUP III.—NITRATE GROUP.

508. This Group includes nitrate, chlorate, bromate and iodate.

The nitrates and chlorates resemble one another in many respects, nore particularly in the fact that none of them are perfectly insoluble a water; hence no method of precipitating them is known.

The bromates and iodates are of comparatively rare occurrence, and only a few of their reactions are therefore described.

NITRATE, (NO₃)'.—Use KNO₃ and Pb(NO₃)₂.

509. FeSO₄-solution gives a dark brown colour with Nitric acid: the nitric acid is formed from the nitrate by the addition of H₂SO₄, and is changed by the action of FeSO₄ into nitrite, which then colours the FeSO₄ (503).

Add to some solution of nitrate, contained in a rather broad test-tube, about an equal volume of strong H₂SO₄. Mix the liquids well, and then cool the mixture by holding the tube in a stream of cold water.

Now pour some strong cold FeSO₄-solution in a gentle stream down the inside of the tube, which is held in an inclined position. A distinct layer of the FeSO₄-solution may thus with care be formed on the surface of the acid liquid. A *dark coloured* film or ring, which sometimes extends upwards, will appear at the surface of contact of the two liquids.

This coloured film is most easily seen when a piece of filter-paper is held between the tube and a bright flame or window, and the light is viewed through the tube and the paper. A mere yellow colour is often produced in the absence of a nitrate, and may be disregarded.

The test is made more delicate by cooling the liquid, since heat prevents or removes the dark colour. For this purpose the heating of the liquid by the mixture of the strong acid with the solution must be avoided as much as possible, and the tube may be allowed to stand in a beaker of cold water for a few minutes.

If much nitric acid is present, the dark colour often extends upwards as soon as the strong sulphuric acid is mixed with the FeSO₄-solution; but the heat which is evolved by the dilution of the strong acid soon causes the NO to escape and to form a reddish-brown gas in the test-tube, and the liquid at the same time loses its dark colour.

510. Copper: when a few small scraps of Cu are dropped into some nitrate solution, and strong H₂SO₄ is added, reddish-brown gas will appear either at once or when the tube is warmed. If the colour is faint, it is best seen by looking down the tube at a white surface.

The liquid at the same time becomes blue from the production in it of copper nitrate:—

$$8HNO_3 + 3Cu = 2NO + 3Cu(NO_3)_2 + 4H_2O.$$

The NO which is evolved is a colourless gas, but it yields reddishbrown NO₂ by oxidation when it mixes with the air in the upper part of the vessel.

511. Indigo solution, if it is poured in a cold and dilute condition into cold nitrate solution, until the mixture becomes faintly but decidedly blue, retains its blue colour unchanged when SO₂ is passed into the liquid (*Note*, 519), or when freshly made H₂SO₃ solution is added. This reaction distinguishes a nitrate from a chlorate (519).

- 512. Brucine: if a little brucine is moistened on a watch-glass with a drop of strong $\rm H_2SO_4$, an intense red colour is produced on adding a drop of solution of a nitrate. This reaction is very delicate.
- 513. Reduction to Nitrite by Ignition: ignition converts alkaline nitrate into nitrite: hence if solid nitrate is heated to redness with fusion mixture $(Na_2CO_3 + K_2CO_3)$ on platinum foil for several minutes, a soluble alkaline nitrite is formed, which may be dissolved when it is cool by boiling the foil in water.

A portion of this solution may be proved to contain nitrite by the starch test (504).

Another portion may be acidified with HNO₃, and will be found to give no precipitate with AgNO₃. Compare with paragraph 521.

514. Reduction to Nitrite by the action of Hydrogen: nitric acid may also be reduced to nitrous acid by the action of nascent hydrogen.

Add dilute H₂SO₄ to some nitrate solution, and then add some freshly-made starch-solution containing KI: no colour will be produced, but when a fragment of Zn is dropped in, and hydrogen is evolved, the liquid will assume a deep blue colour (504).

- 515. Fusion of a Nitrate on Charcoal in the blowpipe-flame causes "deflagration"; that is to say, the surface of the charcoal burns rapidly with a slight hissing noise.
- 516. Ignition of the Nitrate of a Heavy Metal, in a glass tube closed at one end, causes the evolution of a reddish-brown gas with a characteristic smell (NO₂) and of oxygen:—PbN₂O₆ = 2NO₂ + O + PbO. The NO₂ is recognised by its colour and by its smell, and the oxygen is found by inflaming a glowing splinter of wood. Lead nitrate may be used in trying this experiment.
- 517. Hydrogen nitrate, or Nitric acid, HNO_3 , has a strongly acid reaction. It gives with $FeSO_4$ -solution a brown ring (509), and with Cu a reddish-brown gas and a blue solution (510).

When nitric acid is evaporated with quill-clippings it stains them bright yellow.

CHLORATE, (ClO₃)'.—Use KClO₃.

519. Indigo solution, if it is added in a cold dilute condition to a cold solution of chlorate until the liquid is coloured faintly but distinctly blue, loses its colour as soon as SO₂ is passed into the liquid, or when some freshly-made H₂SO₃-solution is added and the mixture is shaken. Difference from nitrate (511).

Note.—Since the presence of sulphuric acid must be avoided in this reaction, it is well to pass SO_2 gas into the liquid which is being tested. The gas may be most readily obtained from a bottle of liquid SO_2 .

520. Strong H₂SO₄, when it is poured upon a little solid chlorate, becomes orange-red in colour, and when the liquid is shaken it evolves a bright yellow gas, ClO₂:—

$$3 \text{KClO}_3 + \text{H}_2 \text{SO}_4 = 2 \text{ClO}_2 + \text{KClO}_4 + \text{K}_2 \text{SO}_4 + \text{H}_2 \text{O}.$$

This gas has a smell somewhat resembling that of chlorine. It undergoes explosive decomposition when it is gently heated, and the acid mixture accordingly crackles and even detonates when it is warmed in the tube by the flame.

Caution.—This experiment may be dangerous, unless very little chlorate is used, and the mixture is then heated gently with the mouth of the test-tube turned away from the person.

521. Ignition of Alkaline Chlorate evolves Oxygen and produces Chloride.

AgNO₃-solution gives no precipitate with solution of a chlorate, if chloride is absent.

But if some of the solid potassium chlorate is heated in a testtube, it will fuse and give off oxygen with effervescence, and the chlorate will be converted into chloride:—

$$\text{KClO}_3 = \text{KCl} + 30.$$

The oxygen may be detected by holding in the tube a slip of wood with a spark at its end; the spark will burst into a flame.

If the tube is then allowed to cool, and the residue is dissolved by heating it with water, a portion of the solution will yield no evidence of the presence of a nitrite (504).

Another portion of the solution will yield a white precipitate (AgCl) on the addition of AgNO₃; this precipitate will not dissolve in HNO₃ even on boiling, but will dissolve readily in AmOH (539): it is therefore produced by the chloride which has been formed from the chlorate, and indirectly proves the presence of chlorate.

- 512. Brucine: if a little brucine is moistened on a watchglass with a drop of strong H₂SO₄, an intense red colour is produced on adding a drop of solution of a nitrate. This reaction is very delicate.
- 513. Reduction to Nitrite by Ignition: ignition converts alkaline nitrate into nitrite: hence if solid nitrate is heated to redness with fusion mixture $(Na_2CO_3 + K_2CO_3)$ on platinum foil for several minutes, a soluble alkaline nitrite is formed, which may be dissolved when it is cool by boiling the foil in water.

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bright yellow.

CHLORATE, (ClO₃)'.—Use KClO₃.

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This gas has a smell somewhat resembling that of chlorine. It undergoes explosive decomposition when it is gently heated, and the acid mixture accordingly crackles and even detonates when it is warmed in the tube by the flame.

Caution.—This experiment may be dangerous, unless very little chlorate is used, and the mixture is then heated gently with the mouth of the test-tube turned away from the person.

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But if some of the solid potassium chlorate is heated in a testtube, it will fuse and give off oxygen with effervescence, and the chlorate will be converted into chloride:—

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The oxygen may be detected by holding in the tube a slip of wood with a spark at its end; the spark will burst into a flame.

If the tube is then allowed to cool, and the residue is dissolved by heating it with water, a portion of the solution will yield no evidence of the presence of a nitrite (504).

Another portion of the solution will yield a white precipitate (AgCl) on the addition of AgNO₃; this precipitate will not dissolve in HNO₃ even on boiling, but will dissolve readily in AmOH (539): it is therefore produced by the chloride which has been formed from the chlorate, and indirectly proves the presence of chlorate.

A chlorate which does not yield a soluble chloride by ignition must be mixed with fusion-mixture before it is ignited for this test.

This reaction readily distinguishes a chlorate from a nitrate (513).

- * 522. When a Chlorate is heated on Charcoal in the blow-pipe-flame it causes the charcoal to deflagrate.
- 523. Hydrogen chlorate, or Chloric acid, HClO₃, is an odourless liquid, which first reddens and then bleaches litmus-paper. When chloric acid is kept for some time, it changes into HClO₄, O, Cl, and H₂O. The same change occurs rapidly when the acid is heated.

DETECTION OF NITRATE AND CHLORATE.

524. There is little difficulty in distinguishing nitrate from chlorate when they occur separately.

But nitrate and chlorate, when they are present together, more or less interfere with one another's reactions. They may be detected in the absence, or after the removal of chloride and nitrite, by heating the solid substance strongly for a short time, and then testing for nitrite and chloride (513, 521).

The preliminary removal of nitrite may be effected by boiling the substance with excess of acetic acid; and chloride may be removed by precipitation with AgNO₃, which must, however, not be added in excess. It is evident that the portion of the solution, to which AgNO₃ has been added, cannot be used for the test for nitrate, but if Ag₂SO₄ is substituted for AgNO₃ this difficulty is avoided.

A method for detecting mixed nitrate and chlorate by the action of nascent hydrogen, produced by zinc-dust in the presence of alkaline hydroxide, and conversion into NH₃ and chloride, is given in paragraph 1287.

DETECTION OF NITRATE IN PRESENCE OF NITRITE.

525. Add to the solution about one gram of urea, and then slowly add dilute sulphuric acid; the nitrous acid will be destroyed:— $CO(NH_2)_2 + 2HNO_2 = 2N_2 + CO_2 + 3H_2O$. When the evolution of gas has ceased, ascertain if any nitrite remains in the solution by adding solution of KI and starch (504), and if necessary add more urea and heat until the complete destruction of nitrite is proved by the absence of blue coloration of the starch.

Then add a little metallic zinc; a blue coloration will now be produced if nitrate is present (514).

The solution of nitrite and nitrate may also be freed from nitrite by boiling it for several minutes with strong ammonium chloride solution:—NaNO₂ + NH₄Cl = NaCl + 2H₂O + N. The solution may then be examined for nitrate as is directed above.

Bromate, (BrO₃)'.—Use KBrO₃.

526. $AgNO_3$: white crystalline precipitate, $AgBrO_3$, almost insoluble in HNO_3 , but readily soluble in AmOH.

527. $\mathbf{Hg}_2(\mathbf{NO}_3)_2$: white precipitate of $\mathbf{Hg}_2(\mathbf{BrO}_3)_2$.

528, $\mathbf{H}_2\mathbf{S}$, or $\mathbf{H}_2\mathbf{SO}_3$: each of these reduces bromate to bromide:— $\mathbf{KBrO}_0 + 3\mathbf{H}_2\mathbf{S} = \mathbf{KR}_2 + 3\mathbf{H}_2\mathbf{O} + \mathbf{S}$

 $\frac{\mathrm{KBrO_3}}{\mathrm{KBrO_3}} + 3\mathrm{H_2S} = \mathrm{KBr} + 3\mathrm{H_2O} + \mathrm{S} + 3\mathrm{H_2SO_3} = \mathrm{KBr} + 3\mathrm{H_2SO_4}.$

529. Mineral acids liberate bromic acid, which then decomposes with formation of Br, H₂O and O; or if a reducing agent is present HBr is formed, and this reacting with unchanged HBrO₃ liberates Br:—

$${\rm HBrO_3} + 5{\rm HBr} = 3{\rm Br_2} + 3{\rm H_2O}.$$

530. Dry reactions: an alkaline bromate is converted by ignition into bromide and oxygen (521): other bromates are decomposed with evolution of bromine and oxygen.

When a bromate is heated on charcoal in the blowpipe-flame it causes deflagration.

IODATE, (IO₃)'.—Use KIO₃.

- 531. AgNO₃: white curdy precipitate, AgIO₃, readily soluble in AmOH, but soluble with difficulty in HNO₃.
 - 532. BaCl₂: white precipitate, Ba $(IO_3)_2$, soluble with difficulty in dilute HCl and dilute HNO₃.
 - 533. $\mathbf{Hg}_2(\mathbf{NO}_3)_2$: light yellow curdy precipitate, $\mathbf{Hg}_2(\mathbf{IO}_3)_2$.
 - 534. H₂S, H₂SO₃: each of these reduces iodate to iodide (528).
 - 535. Dilute H_2SO_4 does not decompose iodate, but if reducing substances, such as H_2SO_3 or $FeSO_4$, are present, HI is produced and reacts with unchanged HIO₃, causing separation of iodine:—HIO₃ + 5HI = $3I_2 + 3H_2O$.
 - 536. Dry reactions: all iodates are decomposed by ignition; some of them lose O and are converted into iodides, others evolve both O and I. When an iodate is heated on charcoal it causes deflagration.

DETECTION OF BROMATE AND IODATE.

537. Bromates and iodates are not of common occurrence. They may be recognized by treatment with dilute H_2SO_4 in the presence of H_2SO_3 or $FeSO_4$ (529, 535), or they may be converted into bromides or iodides (528, 534) and identified as such (546 and 552 et seq.).

GROUP IV.—CHLORIDE GROUP.

538. This Group includes chloride, bromide and iodide.

These Acid-radicles closely resemble one another in many of their reactions. They are precipitated by AgNO₃ from their solutions, even after HNO₃ has been added in excess. In this respect they differ from all acid-radicles except cyanide, thiocyanate, ferrocyanide and ferricyanide; and these are easily distinguished by other means.

The elements are commonly known as the "halogens," and their salts as "halides" of the metals.

CHLORIDE, (Cl)'.—Use NaCl.

539. AgNO₃, when it is added to the solution of a chloride, gives a pure white precipitate of AgCl.

The precipitate collects into curdy masses when the liquid is heated or shaken: it rapidly turns violet in sunlight, and slowly darkens even in diffused daylight.

Shake up and pour off the liquid with the precipitate into several test-tubes.

Decant the water from one portion and warm the precipitate with HNO₃, it will not dissolve: to another portion of the precipitate add AmOH, it will readily dissolve: AgCl may also be shown to be soluble in KCN-solution and in Na₂S₂O₃-solution.

Warm another portion of the precipitate, from which the liquid has been decanted, with a little strong H₂SO₄; the acid will remain colourless and no violet vapour will be given off as is the case with AgI (552).

- 540. Strong H₂SO₄, when it is added to a solid chloride, causes colourless HCl to be given off: this gas may be recognised by fuming in the air, by reddening moistened blue litmus-paper, and by making a drop of AgNO₃-solution, acidified with HNO₃, milky (474, 1-4).
- 541. MnO₂ and H₂SO₄.—Mix together intimately some finely-powdered chloride and MnO₂, then add strong H₂SO₄ and warm the mixture: chlorine gas will be evolved:—

$$2\text{NaCl} + \text{MnO}_2 + 3\text{H}_2\text{SO}_4 = \text{Cl}_2 + \text{MnSO}_4 + 2\text{NaHSO}_4 + 2\text{H}_2\text{O}.$$

This gas is recognised by its power of bleaching a piece of moistened litmus-paper, which is introduced into the tube on a glass rod.

A very delicate way of making the test is to mix the substances in a small beaker, and then cover this with a watch-glass which bears on its under surface a piece of moistened litmus-paper; the mixture is then warmed and allowed to stand for some time, and the test-paper is occasionally examined in order to see if its colour has disappeared.

Moist starch is not coloured when it is held in the escaping gas: this distinguishes chloride from bromide, when they are examined by this test. Compare with paragraph 547.

Many samples of commercial MnO2 evolve CI when they are heated with

 $\rm H_2SO_4$. Hence unless the $\rm MnO_2$ has been specially prepared by precipitation, or is known to give no Cl when it is heated with $\rm H_2SO_4$, it must be heated with $\rm H_2SO_4$ as long as any bleaching action is produced on moist litmuspaper held in the tube; the substance to be tested may then be added, and $\rm Cl$ may be again tested for while the mixture is being heated.

542. Chromyl chloride test.—An intimate mixture is made of a solid chloride with three or four times as much K₂Cr₂O₇, by rubbing the two substances together to a fine powder in a mortar. This mixture is then heated with strong H₂SO₄ in the tube or flask represented in Figure 56 (474), or in a Clarke's retort (Fig. 57).

Reddish-brown vapour of chromyl chloride, CrO₂Cl₂, will be evolved:—

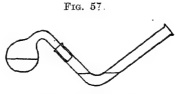
$$\begin{split} 4 \text{NaCl} + \text{K}_2 \text{Cr}_2 \text{O}_7 + 6 \text{H}_2 \text{SO}_4 &= 2 \text{CrO}_2 \text{Cl}_2 + 4 \text{NaHSO}_4 + \\ 2 \text{KHSO}_4 + 3 \text{H}_2 \text{O}. \end{split}$$

If the vapour escaping from the delivery-tube is passed into some water, contained in a test-tube or in the bend of the condenser of Clarke's apparatus (Fig. 57), the water will become reddishyellow in colour, owing to the formation of H_2CrO_4 :—

$$CrO_2Cl_2 + 2H_2O = H_2CrO_4 + 2HCl.$$

When excess of AmOH is added to the reddish-yellow liquid the colour will change to pale yellow, since Am₂CrO₄ is formed.

But the original reddish-yellow colour will be reproduced when excess of HĀ is added; and the presence in this liquid of H₂CrO₄, and therefore of chloride in the original substance, may be proved by the formation of a yellow precipitate (PbCrO₄) on the addition of PbĀ₂-solution.



CLARKE'S RETORT.

Caution.—The detection of a chloride by this method depends on the formation of H₂CrO₄ by the action of the water in the tube on the vapour of CrO₂Cl₂. Great care must therefore be taken that none of the mixture containing chromate or chromic acid is allowed to pass into the delivery-tube and thence into the water, since if this occurred, chromic acid would be found even when no chloride was present.

Care must also be taken that the water is not sucked back into the hot strong H₂SO₄ when the apparatus shown in Fig. 56 is used. This is prevented by raising the delivery-tube out of the water as soon as the heating is stopped.

543. Hydrogen chloride, or Hydrochloric acid, HCl, is a colourless gas, which fumes in the air, and dissolves very readily in water, forming a strongly acid solution. Both the gas and its solution render AgNO₃-solution, which has been acidified with

 HNO_3 , milky. The solution of the acid evolves chlorine when it is heated with MnO_2 , or with other oxidising substances.

544. Chlorine is a greenish-yellow gas with a suffocating odour. It dissolves in about one-third its volume of water, and the "chlorine-water" thus produced is pale yellowish-green in colour and possesses the general properties of the gas.

This solution probably undergoes change to a small extent:— H_2O+Cl_2 —HCl+HClO, and the products ultimately formed, slowly in the dark, more rapidly in day- or sunlight, are HCl and $O:=H_2O+Cl_2=O+2HCl$.

Chlorine readily bleaches many moist vegetable colours, such as indigo and litmus, the bleaching being probably caused by the nascent oxygen produced as is shown by the above equation. Chlorine also liberates iodine from KI-solution, and the liquid becomes yellow or brown in colour.

AgNO₃-solution gives a white precipitate of AgCl with chlorine, but the precipitation is incomplete since soluble silver chlorate is simultaneously formed: $-3\text{Cl}_2 + 6\text{AgNO}_3 + 3\text{H}_2\text{O} = 5\text{AgCl} + \text{AgClO}_3 + 6\text{HNO}_3$.

545. Chlorine and Hydrochloric acid may be detected when they may be present together in solution as follows. After Cl has been found by its characteristic smell and bleaching power, the liquid is shaken up with mercury until the smell of Cl disappears and the liquid no longer bleaches litmus: insoluble Hg₂Cl₂ is formed and the liquid is therefore left neutral in reaction, if HCl was absent: an acid reaction of the liquid after the removal of chlorine therefore indicates the presence of an acid in the original solution, and hydrochloric acid may then be recognised by its characteristic tests (543).

Bromide, (Br)'—Use NaBr.

546. AgNO₃, when it is added to solution of a bromide, gives a y-ellowish-white precipitate (AgBr).

This precipitate is easily coagulated when the liquid containing it is heated or shaken; it is insoluble in HNO₃, easily soluble in KCN-solution and in Na₂S₂O₃-solution, and slightly soluble in AmOH. It resembles the precipitate of AgCl in these respects, but it is distinguished from AgCl by its yellowish colour, and by its slight solubility in AmOH.

If the liquid is decanted and the precipitate is heated with strong H_2SO_4 , no yielet vapour will be evolved as is the case with AgI (552).

547. Strong H₂SO₄, if it is heated with a bromide, causes the evolution of pungent acid fumes, HBr, which are accompanied and coloured by *reddish-brown* vapour of Br. This vapour somewhat resembles Cl in smell and by bleaching litmus, but it differs from Cl in colour.

The bromine vapour may also be recognised by its power of staining cold moist starch orange-red. The starch-powder may be

taken up on the wetted end of a glass rod, and then moistened by breathing upon it several times.

If MnO₂ is mixed with the bromide before H₂SO₄ is added, Br₂ is evolved in greater quantity (541).

548. Chlorine-water, or solution of bleaching-powder which has been acidified with HCl, if it is added carefully drop by drop to solution of a bromide, liberates Br and this colours the solution orange-red.

The addition of Cl in excess must be carefully avoided, since Cl combines with Br and destroys its colour.

Warm a part of the coloured solution; reddish-brown Br-vapour will be given off, and will stain cold moist starch orange-red (547).

Add carbon bisulphide (CS₂) to another portion of the coloured solution, and mix the liquids well by shaking them; the Br will be dissolved away from the water by the CS₂, and if the liquids are allowed to stand at rest for a short time, the reddish-brown solution will sink beneath the colourless water.

If a little KOH-solution is now added and the liquids are shaken up together, the colour of the Br will disappear from the CS₂; this is due to the formation of the colourless salts KBr and KBrO₃, which remain in solution in the water:—

$$3Br_2 + 6KOH = 5KBr + KBrO_3 + 3H_2O.$$

549. Evolution of Bromine on addition of H₂SO₄ and Chromate.—If an intimate mixture of a solid bromide and K₂Cr₂O₇ is heated with strong H₂SO₄ in the apparatus shown in Fig. 57, page 147, reddish-brown bromine vapour will be evolved:—

$$\begin{split} 6 \text{NaBr} + \text{K}_2 \text{Cr}_2 \text{O}_7 + 11 \text{H}_2 \text{SO}_4 &= 3 \text{Br}_2 + \text{Cr}_2 (\text{SO}_4)_3 + 6 \text{NaHSO}_4 \\ &+ 2 \text{KHSO}_4 + 7 \text{H}_2 \text{O}. \end{split}$$

The Br-vapour may be passed into water, and it will colour the liquid reddish-brown.

If CS₂ is then shaken up with part of this brown solution, it will dissolve away the Br and will settle beneath the colourless water as a brown layer.

If another part of the brown aqueous solution of Br is shaken with AmOH or KOH, the colour of the solution will be destroyed.

These reactions serve to detect a bromide in the presence of a chloride. Compare with paragraph 542.

550. Hydrogen bromide, or Hydrobromic acid, HBr, is a colourless, soluble gas, which fumes in the air and closely resembles HCl gas. Its aqueous solution differs from that of HCl by evolving coloured Br-vapour when it is heated with MnO₂.

551. Bromine is a heavy brown liquid which emits suffocating reddishbrown vapour. Cold water will dissolve two and a half per cent. of Br, producing a reddish-brown solution known as "bromine-water." Br is readily identified by its smell, its colour, and its bleaching action, also by its solubility in CS₂, producing a deep reddish-brown liquid.

The detection of HBr and Br when they may be present together may

be effected by a similar procedure to that described in paragraph 545.

IODIDE, (I)'.—Use KI.

552. AgNO₃, when it is added to solution of an iodide, gives

a yellow precipitate, AgI.

This precipitate is easily coagulated when the liquid containing it is heated or shaken: it is insoluble in HNO₃, very slightly soluble in AmOH, but easily soluble in KCN-solution and in Na₂S₂O₃-solution.

The precipitate of AgI differs from the precipitates of AgCI and AgBr by its colour, and by its very slight solubility in AmOH; but it is distinguished with greater certainty by its behaviour when it is heated with strong H₂SO₄ as follows.

If the liquid is decanted from the precipitate, and the precipitate is then heated with strong H₂SO₄, the acid will be coloured red, and violet vapour of iodine will be seen on looking down the test-tube: the coloured vapour is most distinctly seen after the tube has been allowed to cool for a short time.

If a mere trace of iodine vapour is evolved, it may be detected by holding in the test-tube a glass rod which has been dipped into freshly-made starch solution (*Note* 1); the starch will be coloured blue (*Note* 2).

Note 1.—Starch solution is made by stirring a little starch powder with some water in a small porcelain dish, then boiling for a few seconds, and cooling the solution.

Note 2.—The blue colour produced by the action of iodine on starch is commonly stated to be due to the formation of a compound known as "iodide of starch."

The production of this colour requires the presence of an iodide with the starch and iodine, hence the addition of a little KI-solution to the starch-solution makes the above test for iodine vapour more delicate. But the starch solution containing iodide must not be used when substances are present which would liberate iodine from the iodide, and must not be added to a liquid where the introduction of iodide is not permissible.

553. CuSO₄-solution produces a white precipitate of cuprous iodide, Cu₂I₂, and causes the separation of iodine which colours the liquid brown:—

$$4KI + 2CuSO_4 = Cu_2I_2 + I_2 + 2K_2SO_4$$

If the copper sulphate solution has been mixed with solution

of H₂SO₃ or FeSO₄ before it is added to the iodide, the precipitate and liquid are not discoloured by free iodine:—

$$\begin{array}{l} 2 {\rm CuSO_4} + 4 {\rm KI} + {\rm H_2SO_3} + {\rm H_2O} = {\rm Cu_2I_2} + 2 {\rm K_2SO_4} + {\rm H_2SO_4} \\ + 2 {\rm HI}. \end{array}$$

The formation of the precipitate is accelerated by warming the liquid.

Separate this precipitate from the liquid by decantation or by filtration, and warm it with strong $\rm H_2SO_4$; the acid will become red, and violet iodine vapour will be evolved which colours starch solution (552).

Solutions of chloride and of bromide differ from solution of iodide by not being precipitated by CuSO₄-solution.

554. Strong H₂SO₄, if it is warmed with a solid iodide, causes the evolution of acid fumes (HI), accompanied by violet iodine vapour. If the iodine vapour is large in quantity, it will condense on the inside of the test-tube as a black solid.

The iodine vapour colours starch-solution blue. This may be proved by holding in the vapour a glass rod, or a strip of paper, which has been moistened with freshly-made starch-solution (*Notes*, **552**); or by pouring out the heavy iodine vapour into a white porcelain dish, the inside of which has been wetted with the starch-solution.

Iodine is evolved in larger quantity, if MnO_2 has been mixed with the iodide before it is warmed with H_2SO_4 (541).

555. Chlorine-water, or solution of bleaching-powder which has been acidified with HCl, if it is added carefully drop by drop to solution of an iodide sets iodine free. The iodine dissolves in the water colouring it brown: some of the iodine may also separate as a black powder.

The addition of CI in excess must be carefully avoided, as it would form colourless iodine chloride, which does not give the following results.

Divide the brown liquid into three parts in three test-tubes, and proceed as follows:—

- 1. Heat one part; violet iodine vapour will be given off, and will be seen on looking down the test-tube at a white surface. The colour is most distinctly seen if some strong $\rm H_2SO_4$ is added to the liquid before it is heated. If a glass rod is dipped into starch-solution (*Notes*, 552) and is then held in the iodine vapour, the starch will become blue.
- 2. Starch-solution (Note 1, 552), if it is added to another portion of the brown solution, produces an intense blue colour. The

colour often appears black, unless much starch-solution is added and the liquid is largely diluted. When the blue liquid is heated it becomes colourless, but it often regains its colour on being cooled: the liquid must be diluted, and the starch must be added in excess, in order to ensure the removal of the colour by heat.

3. A drop of CS₂ is added to the third portion of the brown solution, and it is well shaken; the CS₂ will dissolve the iodine, and this solution will gradually settle down as a violet layer beneath the water, when the liquids are allowed to stand at rest.

A little KOH solution is then added, and the liquids are shaken together; the violet colour of the CS_2 will disappear, owing to the formation of the colourless salts KI and KIO₃.

556. Nitrous acid, HNO₂.—If a little KNO₂ is dissolved in solution, of an iodide and a dilute acid is added, HNO₂ is produced and liberates the iodine.

A solution of NO_2 in dilute H_2SO_4 (1399) also liberates iodine from an iodide.

The iodine thus liberated may be identified by the three methods described in paragraph 555; of these the starch-test and the CS₂-test are the most delicate.

When either of the two reagents mentioned above is used for liberating iodine from an iodide, it presents the advantage over Cl (555) that it does not hinder the detection of the iodine, even when it has been added in excess. These two reagents also present the advantage that they do not liberate Br from a bromide.

- 557. HgCl₂-solution gives with solution of an iodide a scarlet precipitate, HgI₂, which is easily soluble in excess of solution of HgCl₂ or of the iodide. With a solution of bromide or chloride, HgCl₂ gives no precipitate.
- 558. PbĀ₂-solution gives with solution of iodide a bright yellow precipitate, Pbl₂: if this precipitate is dissolved in the least possible quantity of boiling water, it separates again in the form of golden yellow crystals when the liquid is cooled.

With solution of bromide or chloride, $Pb\tilde{A}_2$ -solution gives a white precipitate.

- 559. Hydrogen iodide, or Hydriodic acid, HI, is a colourless, fuming, soluble gas, resembling HCl and HBr. Its solution differs from solution of HCl or of HBr by evolving iodine when it is heated with MnO₂.
- 560. Iodine is a bluish-black solid which fuses at 114°C. and boils at 184°C., passing into purple vapour. It is soluble in water to the extent of only 0.02 per cent. forming a brown solution, but is more soluble in KI-solution, in alcohol and in other organic liquids. Its solution in most liquids is brown in colour, but the carbon disulphide and chloroform solutions are purple.

Lodine possesses only feeble oxidising power.

It is usually identified by the colour of its vapour, confirmed by the coloration of starch solution (Notes, 552).

DETECTION OF CHLORIDE, BROMIDE, OR IODIDE, WHEN IT OCCURS ALONE.

561. Chloride, Bromide, or Iodide is easily detected by the foregoing tests.

The reactions given in paragraphs 541 and 542 are perhaps the most characteristic tests for a chloride; and the tests described in paragraphs 547-549 serve to distinguish a bromide; an iodide is detected with the greatest ease and certainty by the reaction described in 556.

The colour of the precipitate which has been produced by adding AgNO₃ to the solution made acid with HNO₃, and the behaviour of this precipitate with AmOH, are less trustworthy indications than the above tests; but they often serve to demonstrate which of these acid-radicles is present, since AgCl is *pure white* and is very easily soluble in AmOH, AgBr is *pale yellow* and is not readily soluble in AmOH, while AgI is *primrose-yellow* and is almost insoluble in AmOH.

The action of hot strong H₂SO₄ on the precipitate of AgI, which is described in paragraph 552, is also distinctive of iodide.

DEFECTION OF CHLORIDE, BROMIDE, AND IODIDE, WHEN THEY OCCUR TOGETHER.

562. The detection of these acid-radicles, when they occur together in solution, may be effected by any of the methods described in paragraphs 564, 565, 566.

If chloride is known to be absent, bromide and iodide may be rapidly tested for by the gradual addition of Cl-water in the presence of CS₂ (564), or by fractional precipitation with AgNO₃-solution (566).

563. Insoluble Compounds must be decomposed by means of Na_2CO_3 , in order to obtain the chloride, bromide, and iodide in solution.

Boiling with Na₂CO₃-solution will effect this change in the case of some substances, but heating the finely-powdered substance with three times its weight of fusion mixture (Na₂CO₃ + K₂CO₃) on platinum foil or in a platinum capsule is a more certain and general method.

The embonates are kept in the fused condition with the substance for about ten minutes; the mass is then allowed to cool, and is boiled with water, and the filtered solution is examined by one of the following methods (564-566).

This treatment with fused alkaline carbonates may also be employed for rendering the precipitate which has been produced by AgNO₃ (539, 546, 552) suitable for the following examination.

564,

FIRST METHOD.

564. The most convenient method of examining for chloride, bromide, and iodide, when they may be present together in an aqueous solution, consists in testing for bromide and iodide in one part of the liquid by the chlorine-test (548, 555); then driving off bromine and iodine, if they are present, from another part of the liquid by means of an oxidising solution, and testing for chloride by AgNO₃ (539).

Note.—The aqueous solution of the substance to be tested is referred to below as the "original solution." If the substance is insoluble in water, it must be treated as is directed in paragraph 563, in order to obtain the acid-radicles in solution in water.

To a portion of the original solution, which is to be tested, add a few drops of CS₂: then add to it cautiously, drop by drop, either chlorine-water or acidified solution of bleaching-powder, and shake the liquid after each addition.

If the CS₂ does not become Coloured, bromide and iodide are absent; chloride may then be tested for, by adding to another part of the original solution HNO₃ in excess and AgNO₃ (539).

If the CS_2 becomes Coloured, note the colour which first appears. If the CS_2 becomes brown at once, bromide is present and iodide is absent; if the CS_2 is coloured violet iodide is present, and the cautious addition of the chlorine solution must be continued, constantly shaking the liquid and watching whether, as the violet colour disappears a brown colour takes its place; if the CS_2 turns brown, the presence of bromide is proved: if no brown colour appears, bromide is absent.

Proceed to test for chloride as is directed below.

Remove bromine and iodine from another portion of the original solution by acidifying with dilute H₂SO₄, adding a little potassium permanganate solution (see *Note*), and boiling the solution in a porcelain dish until the brown colour of the liquid disappears. Then add more permanganate solution cautiously and boil the liquid, and continue these processes until the addition of permanganate no longer produces any brown colour, and the liquid retains a pale pink colour for a few seconds: this proves that all bromide and iodide have been removed.

. Now add $AgNO_3$: the formation of a white precipitate will prove the presence of chloride.

Note.—Instead of permanganate one of the following oxidising solutions may be used. (1) Dilute HNO₃, prepared by mixing one volume of acid (142 sp. gr.) with three volumes of water; or (2) cold solution of ammonium persulphate. This must be added in successive quantities, and the liquid must be heated for a time after each addition, until a brown coloration of

the solution is no longer seen. The bromine and iodine will then have been completely removed.

SECOND METHOD.

565. After testing for iodide and bromide by repeatedly shaking part of the original solution (*Note*, 564) with CS₂ during the gradual addition of Clwater, or of acidified CaCl₂O-solution (564), the presence of chloride may be detected, and the presence of iodide and bromide confirmed by successive fractional precipitation with AgNO₃-solution.

For this purpose the rest of the original solution is mixed with AgNO₃-solution (1:100), added drop by drop: yellow AgI will first be formed, and may be removed by shaking or boiling the liquid and filtering off the coagulated AgI: when the yellow AgI is no longer produced and removed, pale-yellow AgBr will appear; and after this has been removed in a similar manner, pure white AgCl will be precipitated. The colour of the silver salt is most easily judged by contrast with white filter-paper.

THIRD METHOD.

566. This method of examination is varied according as iodide is present or absent. Since chloride cannot be detected in the presence of iodide, the iodide must be separated before proceeding to examine for the chloride.

To a small portion of the Original Solution (Note, 564), which has been made just acid by the addition of dilute H_2SO_4 , add a little cold freshly-prepared starch-solution (Note 1, 552). Then add, drop by drop, either strong HNO₃, or solution of nitrous acid (556): the appearance of a dark blue coloration will show the **Presence of Iodide**.

Examine the remainder of the original solution for chloride and bromide, according to the directions given under (a) or (b), the selection of the method depending upon whether iodide has been found or not.

(a) Iodide is Present.—Add to the original solution a mixture of CuSO₄-solution with half its measure of strong H₂SO₃-solution, and warm gently

for a short time: Cu₂I₂ will be precipitated (553).

In order to ascertain whether the iodide has been entirely precipitated, filter a small quantity of the liquid and warm it with a little more of the above reagent. If any further precipitate is produced, return the filtered portion to the unfiltered liquid. Then add more of the reagent to the whole of the liquid, warm it again for a time and test it again, in order to see if the removal of iodide is complete. Repeat these trials until a small portion of the filtrate gives no further precipitate, when it is warmed with more of the reagent.

Then filter, repeating the filtration if necessary, until the liquid is perfectly clear, add KOH in excess to the filtrate, and boil the liquid. Filter off the precipitate thus produced; and test the filtrate, which is now free

from iodide, for bromide and chloride as is directed below (b).

(b) Iodide is Absent.—Make the solution alkaline, if it is not already so, by the addition of pure KOH, and evaporate it to dryness in a porcelain dish. Scrape the residue off the sides of the dish, and mix it with three or four times as much powdered K₂Cr₂O₇ by rubbing the two substances together in the dish with a pestle.

Transfer this mixture to the tube or flask shown in Fig. 56, p. 135 or to a Clarke's retort (Fig. 57, p. 147), taking the greatest care to let none of the

powder remain in the neck or in the delivery-tube.

Then pour in sufficient strong H₂SO₄ to cover the powder to the depth of about a quarter of an inch, and mix the acid and powder by cautious shaking.

Dip the end of the delivery-tube into water contained in a test-tube or receiver and warm the mixture gently, carefully guarding against any of the mixture passing over into the water in the test-tube or receiver.

As soon as no more reddish-brown vapour is visible in the delivery-tube discontinue the heating, and immediately withdraw the tube containing the

water.

If no coloured vapour has been evolved and the water in the tube is colour-

less, chloride and bromide are absent.

If the water is coloured, pour into it sufficient CS₂ to form a layer about half an inch in depth at the bottom of the test-tube. Close the mouth of the tube with the thumb, shake the tube vigorously, and then allow it to stand at rest. If the CS₂, as it settles down, shows a brown colour, this indicates the Presence of Br, derived from bromide.

Separate the water from the CS_2 by pouring it through a wet filter; add to the filtrate AmOH in excess, then HÅ in excess and then $Pb\bar{A}_2$ -solution. The formation of a yellow precipitate indicates the presence of chromate, and this proves indirectly that **Chloride was present** (542) in the original substance.

GROUP V.—PHOSPHATE GROUP.

567. This Group includes phosphate, arsenate, phosphite and hypophosphite; the two former are important and resemble one another closely in their reactions.

PHOSPHORIC ACID AND THE PHOSPHATES.

568. Phosphoric oxide, P_2O_5 , unites with water in different proportions to form three phosphoric acids, orthophosphoric acid H_3PO_4 , pyrophosphoric acid $H_4P_2O_7$, and metaphosphoric acid HPO_3 . When orthophosphoric acid is heated to 250° C. water is given off and pyrophosphoric acid remains; and when either of these acids is heated to redness more water is eliminated and metaphosphoric acid remains.

Where either the pyro- or meta-acid is dissolved in water it recombines with water and forms orthophosphoric acid; this takes place slowly in the cold, but rapidly when the water is heated.

In the ordinary course of analysis pyrophosphate and metaphosphate, if they were originally present, have been converted by solution in water into orthophosphate, hence they must be sought for by special methods applied to the original substance. Овтнорноsрнате, $(PO_4)^{\prime\prime\prime}$.—Use Sodium hydrogen phosphate, Na₂HPO₄.12H₂O.

569. MgSO₄-solution, to which some AmCl has been added •and then a little AmOH, gives with solution of phosphate a white crystalline precipitate, MgAmPO₄.6H₂O.

This precipitate is almost insoluble in AmOH, but is easily soluble in acids. If very little phosphate is present, the precipitate often appears only when the liquid has been gently warmed, then well stirred or shaken, and then allowed to stand for some time.

570. FeCl₃-solution, if it is dropped into phosphate solution, to which a little $H\bar{A}$ and $Na\bar{A}$ -solution have been previously added, gives a yellowish-white precipitate, FePO₄.

571. Ammonium molybdate, AmHMoO₄.—When a drop of phosphate-solution is added to a solution of Am₂MoO₄ in HNO₃, which is assumed to contain AmHMoO₄, a yellow precipitate is produced *in the cold*. (Difference from pyro- and metaphosphate.)

The formation of this precipitate is hastened by gently warming and stirring or shaking the liquid; but the precipitate appears only after a time in very dilute solution of a phosphate. (See *Precautions*.)

A part of the precipitate usually adheres firmly to the inside of the tube; this may be removed by solution of KOH, NaOH, or AmOH, in which it is readily soluble. The precipitate is only slightly soluble in inorganic acids, and is practically insoluble in HNO₃.

Precautions.—This is a most delicate test if it is properly performed. It serves to detect the presence of a mere trace of phosphate, when careful attention is paid to the following precautions.

The AmHMoO4-solution must be prepared according to the directions

given in paragraph 1410, Remark 10.

The solution to be tested must not be alkaline to test-paper. It should be made distinctly acid by addition of HNO₃, and then added in small quantity only to some of the AmHMoO₄-solution in a test-tube. More of the test-solution must only be added if no yellow precipitate forms after the mixed solutions have been gently warmed and stirred.

This last precaution is extremely important, since the presence of an excess

of phosphate altogether prevents the formation of the precipitate.

Show that this is the case, by pouring a few drops of AmHMoO₄-solution into some Na₂HPO₄-solution which has been acidified with HNO₃. No precipitate will form even when the liquid is heated and shaken, since the phosphate is present in large quantity in relation to the AmHMoO₄. If a few drops of this liquid mixture, however, are now added to some fresh AmHMoO₄-solution, the precipitate will appear.

It must also be remembered that the presence of much HCl retards or prevents the formation of this precipitate. A solution of the test-substance, acidified with nitric acid alone, should therefore be used if possible. Dip the end of the delivery-tube into water contained in a test-tube or receiver and warm the mixture gently, carefully guarding against any of the mixture passing over into the water in the test-tube or receiver.

As soon as no more reddish-brown vapour is visible in the delivery-tube discontinue the heating, and immediately withdraw the tube containing the water.

If no coloured vapour has been evolved and the water in the tube is colourless, chloride and bromide are absent.

If the water is coloured, pour into it sufficient CS_2 to form a layer about half an inch in depth at the bottom of the test-tube. Close the mouth of the tube with the thumb, shake the tube vigorously, and then allow it to stand at rest. If the CS_2 , as it settles down, shows a brown colour, this indicates the **Presence of Br**, derived from bromide.

Separate the water from the CS_2 by pouring it through a wet filter; add to the filtrate AmOH in excess, then $H\bar{A}$ in excess and then $Pb\bar{A}_2$ -solution. The formation of a yellow precipitate indicates the presence of chromate, and this proves indirectly that **Chloride was present** (542) in the original substance.

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Where either the pyro- or meta-acid is dissolved in water it recombines with water and forms orthophosphoric acid; this takes place slowly in the cold, but rapidly when the water is heated.

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This precipitate is almost insoluble in AmOH, but is easily soluble in acids. If very little phosphate is present, the precipitate often appears only when the liquid has been gently warmed, then well stirred or shaken, and then allowed to stand for some time.

- 570. FeCl₃-solution, if it is dropped into phosphate solution, to which a little HĀ and NaĀ-solution have been previously added, gives a yellowish-white precipitate, FePO₄.
- 571. Ammonium molybdate, AmHMoO₄.—When a drop of phosphate-solution is added to a solution of Am₂MoO₄ in HNO₃, which is assumed to contain AmHMoO₄, a yellow precipitate is produced in the cold. (Difference from pyro- and metaphosphate.)

The formation of this precipitate is hastened by gently warming and stirring or shaking the liquid; but the precipitate appears only after a time in very dilute solution of a phosphate. (See *Precautions*.)

A part of the precipitate usually adheres firmly to the inside of the tube; this may be removed by solution of KOH, NaOH, or AmOH, in which it is readily soluble. The precipitate is only slightly soluble in inorganic acids, and is practically insoluble in HNO_3 .

Precautions.—This is a most delicate test if it is properly performed. It serves to detect the presence of a mere trace of phosphate, when careful attention is paid to the following precautions.

The AmHMoO₄-solution must be prepared according to the directions

given in paragraph 1410, Remark 10.

The solution to be tested must not be alkaline to test-paper. It should be made distinctly acid by addition of HNO₃, and then added in small quantity only to some of the AmHMoO₄-solution in a test-tube. More of the test-solution must only be added if no yellow precipitate forms after the mixed solutions have been gently warmed and stirred.

This last precaution is extremely important, since the presence of an excess

of phosphate altogether prevents the formation of the precipitate.

Show that this is the case, by pouring a few drops of AmHMoO₄-solution into some Na₂HPO₄-solution which has been acidified with HNO₃. No precipitate will form even when the liquid is heated and shaken, since the phosphate is present in large quantity in relation to the AmHMoO₄. If a few drops of this liquid mixture, however, are now added to some fresh AmHMoO₄-solution, the precipitate will appear.

It must also be remembered that the presence of much HCl retards or prevents the formation of this precipitate. A solution of the test-substance, acidified with nitric acid alone, should therefore be used if possible.

- 572. AgNO₃: yellow precipitate, Ag₃PO₄: pour off a portion of the liquid with the precipitate and show that the precipitate is soluble in AmOH and in HNO₃.
- 573. Phosphine test.—A very delicate test for the presence of P, occurring in the form of phosphate or otherwise, consists in strongly heating the dry solid substance with Mg-powder, or with a small piece of Na, in a little tube of hard glass closed at one end.

When the cool tube is broken and the powdered substance is breathed upon, an onion-like smell will be perceived; this is due to the production and escape of PH₃, which has been formed by the action of the moisture of the breath upon the metallic phosphide.

- 574. Flame-coloration.—If a phosphate is made into a paste with strong $\rm H_2SO_4$, and the mixture is then strongly heated at the tip of the inner blowpipe-flame, it gives a *bluish-green* colour to the flame.
- 575. Blowpipe-test.—Many phosphates, when they are moistened on charcoal with cobalt nitrate solution, and are then heated in the outer blow-pipe-flame, become blue.
- 576. Hydrogen orthophosphate, or Orthophosphoric acid, H_3PO_4 , is a colourless crystalline substance. Its solution is strongly acid, and differs from H_2SO_4 by not charring paper which has been dipped into it and then dried by heat. The free acid does not coagulate albumin. The presence of H_3PO_4 may be detected by reactions 569, 571.

Рукогнозрнате, $(P_2O_7)^{iv}$.—Use Sodium pyrophosphate, $Na_4P_2O_7$.

- 577. Sodium pyrophosphate is made by heating solid sodium orthophosphate, Na_2HPO_4 , to redness on platinum-foil or porcelain until water is no longer driven of :— $2Na_2HPO_4 = Na_4P_2O_7 + H_2O$. The cold residue is then dissolved in *cold water*, since the solution of pyrophosphate is gradually converted by heat into orthophosphate, and the solution is used at once for the following reactions.
- 578. AgNO $_3$: heavy white precipitate of silver pyrophosphate, readily soluble in AmOH and in ${\rm HNO}_3$. (Difference from orthophosphate.)
- 579. AmHMoO₄ produces no precipitate in the cold; but the pyrophosphate is converted into the orthophosphate when the liquid is heated, and yellow ammonium phosphomolybdate is precipitated (571).
- 580. MgSO₄; white precipitate, soluble in excess of either solution; but when the mixture is boiled, a precipitate is formed which does not disappear on cooling. (Difference from metaphosphate.)
- 581. Albumin-solution is not coagulated by free pyrophosphoric acid.
 (Difference from metaphosphoric acid.)

METAPHOSPHATE, (PO3)'.—Use Sodium metaphosphate, NaPO3.

582. Sodium metaphosphate is obtained by heating either microcosmic salt or NaH₂PO₄ to redness on platinum foil or porcelain:—NaNH₄HPO₄ = NaPO₃ + NH₂ + H₂O. The cold residue is dissolved in cold water since,

like pyrophosphate, metaphosphate is slowly converted into orthophosphate when its solution is heated: this solution is used at once for the following reactions.

- 583. AgNO₃: white flocculent precipitate of silver metaphosphate, soluble in AmOH and in HNO₃.
- $584.\ \mathrm{MgSO_4}:$ no precipitate from moderately dilute solution of metaphosphate even on boiling the liquid. (Difference from ortho- and pyrophosphates.)
- 585. AmHMoO₄: no precipitate in the cold, but when the mixture is boiled yellow ammonium phospho-molybdate is precipitated (571, 579).
- 586. Albumin-solution is coagulated by metaphosphoric acid. Prove this by acidifying the metaphosphate-solution with HĀ in order to form metaphosphoric acid, and adding this solution to white of egg dissolved in water, the liquid will become white and semi-opaque. (Difference from orthoand pyrophosphoric acids.)

ARSENATE, (AsO₄)".-Use Na₂HAsO₄.12H₂O solution.

587. The precipitates formed in reactions 569–571 by a phosphate are precisely similar in appearance and general properties to those formed by the same reagents with solution of an arsenate; but AmHMoO₄ gives a precipitate with an arsenate only when the liquid is boiled, and not when it is gently heated, as in the case of a phosphate.

The precipitate (Åg₃AsO₄) which AgNO₃ gives with an arsenate is *brown*, while that given by a phosphate is *yellow* (572).

DETECTION AND SEPARATION OF PHOSPHATE, ARSENATE AND ARSENITE.

- 588. The following Differences serve to detect and separate phosphate and arsenate:—
- 1. The fact that an arsenate yields a precipitate with AmHMoO₄ only when the liquid is *boiled*, whereas the corresponding precipitate is produced by a phosphate when the liquid is only *gently heated*.
- 2. Arsenate solution, which has been boiled with strong HCl, gives, when H₂S is passed into the hot liquid, first a white precipitate of S and then yellow As₂S₃; phosphate solution under the same circumstances gives no precipitate.
- 3. In perfectly neutral arsenate solution AgNO₃ gives a brown precipitate; in neutral phosphate solution it gives a cancry yellow precipitate.

DETECTION OF PHOSPHATE OR ARSENATE.

589. The Simplest Method of ascertaining which of these two Acid-radicles is present is to throw the precipitate, which has been produced by the addition of AmCl, AmOH and MgSO4 (569) upon a filter; then wash it with a little cold water, and drop AgNO₃-solution upon it. If the precipitate consists of MgAmAsO₄, it will become brown; if it is MgAmPO4, it will become canaryyellow.

The precipitate, which has been produced by the addition of AmCl, AmOH and MgSO4, may also be examined for arsenate by dissolving part of it in dilute HNO3, adding AgNO3-solution as long as it causes any white precipitate, and then adding very dilute AmOH gradually drop by drop.

If arsenate is present, a brown precipitate will form just before the liquid becomes alkaline to test-paper: a phosphate will give

a yellow precipitate.

Instead of mixing the dilute AmOH with the acid liquid, the AmOH may be gently poured in upon the top of the liquid: the colour will then appear at the surface of contact of the ammonia solution with the acid liquid.

DETECTION OF ARSENATE, PHOSPHATE AND ARSENITE, WHEN THEY ARE PRESENT TOGETHER.

590. A clear mixture of solutions of MgSO₄, AmCl and AmOH is added as long as it causes any further precipitate, the liquid is filtered and the filtrate is preserved.

A small portion of the precipitate is then tested at once for arsenate by dropping $AgN\hat{O}_3$ -solution upon it. If the precipitate turns yellow, Phosphate is present, and Arsenate is absent. If the precipitate turns brown, Arsenate is present, and phosphate may

The arsenate must be separated from the other part of the precipitate before phosphate can be tested for.

The rest of the precipitate is accordingly dissolved in a little boiling strong HCl, and H₂S is passed for several minutes into this hot solution; the presence of an arsenate will be confirmed by the liquid becoming milky owing to the separation of white S, which is followed by the precipitation of yellow As₂S₃ (174).

This precipitate is filtered off and H2S is again passed into the boiling filtrate. If any further precipitate is produced, the gas must be further passed until it no longer causes a precipitate in

The precipitation by H_2S is rendered complete without delay, if the original solution in HCl is first treated with H_2SO_3 . See paragraph 174.

The liquid is then filtered and the clear filtrate is evaporated just to dryness, the residue is dissolved in a little dilute HNO₃, and this solution is tested for phosphate by adding a few drops of it to AmHMoO₄-solution and gently warming the liquid: a yellow precipitate will indicate the **presence of Phosphate** (571).

Arsenite may be tested for by acidifying the filtrate, which has been reserved from the original Mg-precipitate, and passing H₂S into it: yellow As₂S₃ will be precipitated at cance if arsenite is present.

PHOSPHITE, (PHO3)".—Use solution of Na2PHO3.

- 591. AgNO₃: black precipitate, Ag, when the liquid is allowed to stand or is heated. The addition of AmOH in small quantity increases the delicacy of this test, but it must be remembered that the original precipitate is soluble in excess of this reagent.
- 592. HgCl₂: white precipitate, Hg₂Cl₂, which becomes grey, Hg, when the liquid in which it is suspended is heated.
 - 593. PbA2: white precipitate, PbPHO3, which is insoluble in acetic acid.
 - 594. BaCl₂: white amorphous precipitate, soluble in all acids.
- 595. Reduction reactions.—Solution of phosphorous acid, or of a phosphite which has been acidified with HĀ, acts as a reducing agent: but its action is not as powerful as that of hypophosphorous acid (601), as the following reactions will show:—
- (a) A drop of KMnO₄-solution is not reduced and decolorised by phosphorous acid: but reduction, attended with loss of colour, takes place when the liquid is heated.
 - (b) A drop of CuCl₂- or of CuSO₄-solution is not reduced and decolorised even when the liquid is heated more strongly.
- 596. Solid Phosphite, when it is heated, gives off a mixture of hydrogen and hydrogen phosphide: the latter gas is known by its garlic smell: the mixture of gases is spontaneously inflammable in contact with air, and burns with a bright white flame emitting white fumes: red phosphorus is usually deposited in the ignition-tube and sodium phosphate remains.

HYPOPHOSPHITE, (PH2O2)'.—Use solution of NaPH2O2.

- 597. AgNO₃: white precipitate, AgPH₂O₂, which gradually becomes black; the change is more rapid when the liquid is heated, and is due to the formation of metallic silver.
- 598. HgCl₂: white precipitate, Hg₂Cl₂, which forms slowly on standing, but more rapidly when the liquid is heated.
 - **599.** $Pb\overline{A}_2$: no precipitate. (Difference from phosphite.) Q.A.

DETECTION OF PHOSPHATE OR ARSENATE.

589. The Simplest Method of ascertaining which of these two Acid-radicles is present is to throw the precipitate, which has been produced by the addition of AmCl, AmOH and MgSO₄ (569) upon a filter; then wash it with a little cold water, and drop AgNO₃-solution upon it. If the precipitate consists of MgAmAsO₄, it will become brown; if it is MgAmPO₄, it will become canaryyellow.

The precipitate, which has been produced by the addition of AmCl, AmOH and MgSO₄, may also be examined for arsenate by dissolving part of it in dilute HNO₃, adding AgNO₃-solution as long as it causes any white precipitate, and then adding *very dilute* AmOH gradually drop by drop.

If arsenate is present, a *brown* precipitate will form just before the liquid becomes alkaline to test-paper: a phosphate will give a *yellow* precipitate.

Instead of mixing the dilute AmOH with the acid liquid, the AmOH may be gently poured in upon the top of the liquid: the colour will then appear at the surface of contact of the ammonia solution with the acid liquid.

DETECTION OF ARSENATE, PHOSPHATE AND ARSENITE, WHEN THEY ARE PRESENT TOGETHER.

590. A clear mixture of solutions of MgSO₄, AmCl and AmOH is added as long as it causes any further precipitate, the liquid is filtered and the filtrate is preserved.

A small portion of the precipitate is then tested at once for arsenate by dropping AgNO₃-solution upon it. If the precipitate turns yellow, *Phosphate is present*, and *Arsenate is absent*. If the precipitate turns brown, *Arsenate is present*, and phosphate may also be present.

The arsenate must be separated from the other part of the precipitate before phosphate can be tested for.

The rest of the precipitate is accordingly dissolved in a little boiling strong HCl, and H₂S is passed for several minutes into this hot solution; the presence of an arsenate will be confirmed by the liquid becoming milky owing to the separation of white S, which is followed by the precipitation of yellow As₂S₃ (174).

This precipitate is filtered off and H₂S is again passed into the boiling filtrate. If any further precipitate is produced, the gas must be further passed until it no longer causes a precipitate in the boiling liquid.

The precipitation by H_2S is rendered complete without delay, if the original solution in HCl is first treated with H_2SO_3 . See paragraph 174.

The liquid is then filtered and the clear filtrate is evaporated just to dryness, the residue is dissolved in a little dilute HNO₃, and this solution is tested for phosphate by adding a few drops of it to AmHMoO₄-solution and gently warming the liquid: a yellow precipitate will indicate the **presence of Phosphate** (571).

Arsenite may be tested for by acidifying the filtrate, which has been reserved from the original Mg-precipitate, and passing H₂S into it: yellow As₂S₃ will be precipitated at cance if arsenite is present.

PHOSPHITE, (PHO3)".—Use solution of Na2PHO3.

591. AgNO₃: black precipitate, Ag, when the liquid is allowed to stand or is heated. The addition of AmOH in small quantity increases the delicacy of this test, but it must be remembered that the original precipitate is soluble in excess of this reagent.

592. \mathbf{HgCl}_2 : white precipitate, $\mathbf{Hg}_2\mathrm{Cl}_2$, which becomes grey, \mathbf{Hg} , when the liquid in which it is suspended is heated.

593. $\mathbf{Pb\bar{A}}_{2}$: white precipitate, \mathbf{PbPHO}_{3} , which is insoluble in acetic acid.

594. BaCl2: white amorphous precipitate, soluble in all acids.

- 595. Reduction reactions.—Solution of phosphorous acid, or of a phosphite which has been acidified with $H\bar{A}$, acts as a reducing agent: but its action is not as powerful as that of hypophosphorous acid (601), as the following reactions will show:—
- (a) A drop of KMnO₄-solution is not reduced and decolorised by phosphorous acid: but reduction, attended with loss of colour, takes place when the liquid is heated.
 - (b) A drop of CuCl₂- or of CuSO₄-solution is not reduced and decolorised even when the liquid is heated more strongly.
- 596. Solid Phosphite, when it is heated, gives off a mixture of hydrogen and hydrogen phosphide: the latter gas is known by its garlic smell: the mixture of gases is spontaneously inflammable in contact with air, and burns with a bright white flame emitting white fumes: red phosphorus is usually deposited in the ignition-tube and sodium phosphate remains.

Hypophosphite, (PH2O2)'.—Use solution of NaPH2O2.

597. AgNO₃: white precipitate, AgPH₂O₂, which gradually becomes black; the change is more rapid when the liquid is heated, and is due to the formation of metallic silver.

598. HgCl₂: white precipitate, Hg₂Cl₂, which forms slowly on standing, but more rapidly when the liquid is heated.

599. $\mathbf{Pb\bar{A}}_2$: no precipitate. (Difference from phosphite.) Q.A.

600. BaCl₂: no precipitate. (Difference from phosphite.)

- 601. Reduction reactions.—Solution of hypophosphorous acid, or of a hypophosphite which has been acidified with HA, acts as a powerful reducing reagent: this is proved by the following reactions:—
 - (a) A drop of KMnO₄-solution is reduced and decolorised, when it is added to the cold solution.
 - (b) When the solution is mixed with a little CuSO₄-solution and the mixture is heated to 55° C., red copper hydride, CuH₂, is precipitated, and this evolves H and leaves metallic copper when the liquid is heated more strongly.
 - (c) When the solution is mixed with a little CuCl₂-solution, the copper salt is decolorised and white Cu₂Cl₂ is precipitated; the Cu₂Cl₂ then gradually changes to red metallic copper.
- 602. Solid Hypophosphite, when it is heated, gives off hydrogen phosphide; this gas may be recognised by its garlic smell, and by kindling spontaneously in contact with the air: it burns with a bright white flame which emits white fumes and usually deposits red phosphorus on the interior of the ignition tube. The residue in the tube consists of sodium phosphate.

GROUP VI.—PER-SALT GROUP.

603. This Group includes permanganate and the related manganate, persulphate, perchlorate and periodate; also percarbonate and perborate, which are unstable and of less importance.

Hydrogen peroxide, H₂O₂, and ozone are also included in this group.

The above grouping is somewhat indefinite, since it associates the salts of the higher ϵ cids, HMnO₄, HClO₄, HIO₄ and the salts of the true per-acids H₂S₂O₈, H₂C₂O₆ and HBO₄. Many of the rarer per-acids and their salts are not described here.

PERMANGANATE, (MnO₄)'.—Use solution of KMnO₄.

Solid permanganates are usually very dark red or purple in colour; their solutions are usually purple or reddish-purple.

. 604. KOH, if it is added in large excess, causes the purple colour of the solution to change to green: this is due to the conversion of the permanganate into manganate:—

 $2KMnO_4 + 2KHO = 2K_2MnO_4 + H_2O + O.$

The purple colour reappears when this alkaline green liquid is acidified (610).

605. H₂O₂, when it is added to a permanganate solution which has been acidified with H₂SO₄, reduces the permanganate to a manganese salt with evolution of oxygen:—

$$2KMnO_4 + 3H_2SO_4 + 5H_2O_2 = 2MnSO_4 + K_2SO_4 + 8H_2O.$$

- 606. Reducing reagents.—The addition of SO₂, H₂S, acidified solution of nitrite, or of other reducing reagents, at once destroys the purple colour of permanganate solution, and a nearly colourless manganese salt is formed. It follows that in the ordinary course of analysis permanganate will be converted into a manganous salt, which will be detected in Group III.B.
- 607. Heated with Acid.—Solid permanganate, or its strong solution, gives off oxygen when it is heated with dilute H₂SO₄, and chlorine gas when it is heated with HCl: the purple colour of the permanganate at the same time disappears. If the sulphuric acid is strong, purple vapour of Mn₂O₇ is evolved, which causes explosion since it is very unstable.

MANGANATE, (MnO₄)".—Use alkaline solution of K₂MnO₄.

608. Solid manganates are almost black: their solutions are bluish-green in colour.

609. H₂O.—Manganates undergo decomposition in contact with water, with production of purple permanganate and MnO₂, unless much alkaline hydroxide is present:—

$$3K_2MnO_4 + 2H_2O = 2KMnO_4 + 4KOH + MnO_2$$

610. Acid, if it is added in sufficient quantity, causes the green solution of manganate to change to reddish-purple solution of permanganate: even carbon dioxide produces this change, and it may therefore be brought about by exposure of the manganate solution to the air:—

$$3 {\rm K_2MnO_4} + 3 {\rm H_2SO_4} = 2 {\rm KMnO_4} + 2 {\rm K_2SO_4} + {\rm MnSO_4} + 3 {\rm H_2O} + 0.$$

The above change is reversed when solution of an alkaline hydroxide is added in excess to the red solution (604).

611. Reducing reagents decolorise an acidified solution of manganate and produce manganous salt (606).

Persulphate, (S₂O₈)".—Use Ammonium persulphate, Am₂S₂O₈, dissolved in *cold* water.

612. When a strong cold solution is boiled, it gives off oxygen which may be collected and identified by causing a glowing wooden

splinter to burst into flame: the persulphate is meanwhile reduced to sulphate which remains in solution.

- 613. BaCl₂ gives a precipitate only when the solution is boiled. Since, however, persulphate almost invariably contains sulphate as a product of decomposition, the test should be made by adding BaCl₂ in excess, filtering off any BaSO₄ through a double filter paper, and then boiling the clear filtrate: the formation of a white precipitate by boiling indicates the presence of persulphate.
- 614. Mn- or Pb-salts give a brown precipitate of the corresponding hydrated peroxide when the liquid is boiled: Ni- and Co-salts give a similar result in alkaline solution of persulphate.
- 615. KMnO₄, in acidified solution, is not decolorised by persulphate. (Difference from H₂O₂).

PERCHLORATE, (ClO₄)'.—Use KClO₄ solution.

- 616. The perchlorates are more stable than the chlorates. An aqueous solution of perchloric acid is stable and may be preserved indefinitely even when exposed to the light, if it does not contain more than 71 per cent. of the acid. All perchlorates are soluble in water.
- 617. Action of acids.—Perchlorate is not decomposed when it is boiled with HCl. (Difference from chlorate.)
- $\rm H_2SO_4$ has no action on perchlorate in the cold, but the hot strong acid forms $\rm HClO_4$ which may be distilled if proper precautions are observed.
- 618. Potassium salts precipitate from $\mathrm{HClO_4}$ -solution the relatively insoluble, white, crystalline $\mathrm{KClO_4}$; alcohol causes the precipitation to be complete.
- **619.** $\mathbf{H}_2\mathbf{SO}_3$ -solution does not reduce perchlorate (519). (Difference from chlorate.)
- 620. Dry reactions.—Perchlorates require a higher temperature than chlorates for their decomposition by heat, but they deflagrate when they are heated on charcoal in the blowpipe-flame (522).
- **621.** Periodate, Periodic acid.—Periodic acid, $\mathrm{HIO_4}$, is unknown; but para-periodic acid has been prepared. It has the formula $\mathrm{HIO_4.2H_2O}$, or $\mathrm{H_5IO_6}$, and forms deliquescent crystals which fuse at 133° C. and decompose at 150° C. into $\mathrm{I_2O_5}$, $\mathrm{H_2O}$ and O.

Sodium periodate, $Na_2H_3IO_6$, and many complex periodates are also known.

Periodic acid and its salts are more stable than the corresponding chlorine compounds.

622. Percarbonate.—Free percarbonic acid is unknown, but its potassium salt (K₂C₂O₆), which is pale blue in colour, is stable in the dry state.

Moisture decomposes this salt into KHCO₃ and H₂O₂. Cold dilute H₂SO decomposes it with evolution of CO₂ and formation of KHSO₄ and H₂O₂₄.

The salt therefore acts as an oxidising agent like H₂O₂, and gives all the characteristic reactions of this substance.

Percarbonate may be distinguished from H_2O_2 by adding it in small quantity and in a powdered condition to excess of KI-solution; if $K_2C_2O_6$ is present, I will be set free at once:— $K_2C_2O_6+2KI=2K_2CO_3+I_2$; if only KHCO3 and H_2O_2 are present the I is liberated much more slowly.

AgNO₃ and BaCl₂-solutions give with percarbonate white precipitates which are soluble in dilute HNO₃.

623. Perborate.—Sodium perborate, NaBO $_3$.4H $_2$ O, is a white crystalline salt, which is stable in the air. Its aqueous solution is alkaline and contains H_2O_2 : the solution decomposes when it is heated above 40° C. and oxygen is evolved.

Perborate solution liberates I from KI-solution, and oxidises solution of ferrous salt to ferric salt, chromic acid to perchromic acid, and manganous salt to manganese dioxide.

When perborate solution is added to dilute H_2SO_4 containing 50 per cent. of acid, H_3BO_3 is precipitated and H_2O_2 remains in solution.

Sodium perborate solution may be shown to bleach blue litmus readily, and its main industrial application is for bleaching.

Hydrogen Peroxide, H_2O_2 .—Use a 3 per cent., or 10-volume, solution of H_2O_2 .

- **624.** A little H_2O_2 -solution may be prepared by adding a small quantity of sodium peroxide to *ice-cold* water; this will produce a clear strongly alkaline solution which gives all the reactions of H_2O_2 . When this solution is to be used for the "acid reactions" described below it must be acidified with dilute H_2SO_4 , carefully keeping the liquid cool during the addition of the acid.
- 625. Titanium sulphate, $Ti(SO_4)_2$, gives to acid H_2O_2 -solution a distinct yellow colour, which is due to the formation of pertitanic acid:— $TiO_2 + H_2O_2 = TiO_3 + H_2O$. On the addition of an alkali a yellowish orange precipitate is formed which dissolves in excess of the reagent. This is the most delicate test for H_2O_2 . (Difference from persulphate.)
- 626. Chromic acid.—Add to an acid solution of the peroxide a little ether free from alcohol, and a few drops of K₂Cr₂O₇-solution, and shake the liquids; the upper ethereal layer will be coloured *blue* by the formation of perchromic acid. (Difference from persulphate.)
- 627. Reducing action.—KMnO₄ in acid solution is at once decolorised by $\rm H_2O_2$ with the evolution of oxygen:— $\rm 5H_2O_2 + 2KMnO_4 + 3H_2SO_4 = K_2SO_4 + 2MnSO_4 + 8H_2O + 5O_2$. Many oxides, such as Ag₂O, PbO₂, MnO₂, Co₂O₃, are reduced by $\rm H_2O_2$ with evolution of oxygen in a similar way. (Difference from persulphate.)

- 628. Potassium iodide and Starch: if a little starch-paste containing KI-solution (552, Notes 1, 2) is added to the acidified H_2O_2 solution; a blue colour will appear at once:—2KI + H_2O_2 = 2KOH + I_2 .
- 629. $AuCl_3$ is reduced by alkaline H_2O_2 -solution with separation of the metal and evolution of oxygen. The gold separates in a very finely divided state, and appears brown by reflected light and blue by transmitted light.
- 630. Mn- or Co-salt in solution gives a dark coloured precipitate of the hydrated peroxide of the metal. Hypochlorite solution gives the same reaction with Mn- or Co-salt, but it does not reduce gold chloride solution (629).

OZONE, O3.

631. Ozone is always formed where oxygen is exposed to the silent electric discharge. It is also formed to some extent in many reactions which liberate oxygen, as when BaO_2 , MnO_2 or $KMnO_4$ is mixed with sulphuric acid. Ozone is also formed during the slow oxidation of many substances in the air, notably by the oxidation of ordinary phosphorus. It is a strong oxidising agent and behaves like H_2O_2 in many respects.

Difference between Hydrogen Peroxide (H_2O_2) and Ozone in their (O_3) Reactions.

- 632. Ozone and hydrogen peroxide may be distinguished by the following reactions.
 - (a) Iodine is set free immediately by ozone from dilute neutral potassium iodide solution: H₂O₂ requires the presence of free acid for the liberation of I (628).
 - (b) Bromine is liberated by ozone from an acid solution of sodium bromide : this effect is not produced by $\mathbf{H}_2\mathbf{O}_2$.
 - (c) A strip of silver-foil which has been polished with emery-paper and then heated for a moment in the Bunsen-flame is blackened when it is brought into contact with ozone, but not by contact with H₂O₂.
 - (d) No yellow colour is produced by ozone in titanium sulphate solution, but this colour is produced by H₂O₂ (625).
 - (e) Gold is not precipitated from its solution by ozone, but it is precipitated by ${\rm H_2O_2(629)}$.

VII.—UNGROUPED ACID-RADICLES.

633. The following Acid-radicles are not included in any of the preceding Groups:—Borate, Chromate, Silicate, Fluoride, together with the rarer Acid-radicles, Tungstate, Molybdate, Selenite, Tellurate, Tellurate, Titanate and Vanadate.

The four Acid-radicles first mentioned present no marked resemblances to one another or to other Acid-radicles in their reactions: they are therefore usually detected by special tests.

The reactions of the rarer acid-radicles have been already described under the metals, since they are precipitated by Group-reagents for the metals.

BORATE, $(BO_3)'''$.—Use Sodium pyroborate, or Borax, $Na_2B_4O_7.10H_2O$.

634. Turmeric-test.—Powder some borate finely, and stir it with a little dilute HCl on a watch-glass. Immerse in this liquid one-half of a strip of turmeric-paper, and dry the strip on a watch-glass in the steam-oven (48), upon a water-bath (38), or above a small flame.

The part of the test-paper which was moistened will now appear reddish-brown, and will become bluish-black when it is moistened with AmOH.

635. Alcohol-flame Coloration.—Pour alcohol upon some finely-powdered borate in a porcelain dish, add a little strong H₂SO₄, then heat the mixture and kindle the vapour of the spirit.

The flame will show a *green* border, which is due to the formation and combustion of ethyl borate; the colour of the flame is often more distinctly seen when the vapour is relighted after the flame has burnt for a time and has then been extinguished.

This flame-coloration is similar to that which is given by certain other substances; but its spectrum, consisting of four equidistant green lines, is quite characteristic (Fig. 51, p. 50).

636. Blowpipe-flame Coloration.—If a mixture containing CaF₂, KHSO₄ and a borate is finely powdered, then moistened and heated on a loop of platinum wire in the inner blowpipe-flame for a short time, it will colour the outer flame momentarily *green*.

This test is not always decisive, since many phosphates and copper salts

give a similar result.

The flame-coloration is also liable to be masked by the colorations of other substances; but even when this is the case the four equidistant green lines of the boric acid spectrum will be visible, and will afford conclusive evidence of the presence of boric acid.

Most borates also give the green flame coloration and the spectrum when

they are moistened with strong H_2SO_4 and held in the Bunsen-flame on a loop of platinum wire.

- 637. Barium-Calcium- and Lead-salts produce, in fairly concentrated solutions of borate, white precipitates of the metaborates, which are soluble in AmCl-solution.
- 638. Hydrogen borate, or Boric acid (H_3BO_3), usually occurs in lustrous crystalline scales. When this substance is strongly heated, it yields a fusible glass consisting of B_2O_3 .

The acid is soluble both in water and in alcohol; and when these solutions are evaporated, the boric acid partially escapes with the vapour of the liquid. The solution of boric acid turns the colour of blue litmus-paper to wine-red. The reactions in paragraphs 634, 635 and 636 serve to detect boric acid.

CHROMATE, (CrO₄)".—Use Potassium dichromate, K₂Cr₂O₇.

All chromates are more or less yellow or red in colour. The yellow solution of a normal chromate becomes orange-red when an acid is added, owing to the formation of a dichromate: while the reddish solution of a dichromate becomes pale yellow when it is made alkaline, owing to its conversion into normal chromate. See remarks under Chromium, paragraph 281.

639. Reduction of Chromate to Chromic salt.—Various substances, which readily combine with oxygen, deoxidise acidified chromate solution and produce a chromic salt; the colour of the solution at the same time changes from *orange-red* to *bright green*.

Add dilute HCl or H₂SO₄ to some chromate solution, and deoxidise a separate portion of this liquid by each of the following methods: in every case the colour of the liquid will change to bluish-green:—

- (a) Pass H₂S into the hot solution; white S will separate owing to oxidation of the hydrogen of the H₂S.
- (b) Add NaHSO3 or H2SO3 and warm.
- (c) Add alcohol and boil: the smell of aldehyde will be noticed.
- (d) Boil for some time, after adding much strong HCl: Cl gas will be slowly evolved.
- (e) Add Zn and warm: the reducing action of the nascent H will be slow.

. The following equations show the reactions which occur with $\mathrm{H}_2\mathrm{S}$ and with HCl :—

 $K_2Cr_2O_7 + 3H_2S + 8HCl = 2CrCl_3 + 2KCl + 7H_2O + 3S.$ $K_2Cr_2O_7 + 14HCl = 2CrCl_3 + 2KCl + 7H_2O + 3Cl_2.$

The green solution, which results from the above reactions, will give all the reactions for a chromic salt (282-285); and since chromic acid is reduced by boiling HCl and by $\rm H_2S$ to chromic salt, the presence of a chromate will lead to the precipitation and detection of $\rm Cr(OH)_3$ in the ordinary course of analysis. This chromic hydroxide, however, will be known, from the colour of the original solution, to be the result of the reduction of chromate.

640. Hydrogen peroxide, H₂O₂, or Barium peroxide, BaO₂, produces in a cold acidified solution of chromate a characteristic intense but evanescent *blue coloration* of perchromic acid.

This colour is far more permanent if ether is first added, and the liquid is then vigorously shaken immediately after the addition of the peroxide: the ether assumes and retains a blue colour.

Add a few drops of dilute $\rm H_2SO_4$ to 2 c.c. of $\rm H_2O_2$ -solution and shake this liquid with 2 c.c. of ether: then add a little of the chromate solution and shake the liquid again. In the presence of only 0·1 milligramme of chromic acid, the upper ether layer is coloured intensely blue, and the coloration is distinctly visible when only 0·01 milligramme of chromic acid is present.

- 641. PbĀ₂: yellow precipitate, PbCrO₄: soluble in KOH, insoluble in HĀ.
- **642.** $AgNO_3$: purple-red precipitate, Ag_2CrO_4 : soluble in AmOH and in HNO_3 .
- 643. BaCl₂: yellowish-white precipitate, BaCrO₄: insoluble in HĀ, soluble in HCl.
- 644. Hydrogen chromate, or Chromic acid, H₂CrO₄, has not been isolated, but is believed to be present in the reddish-yellow liquid which is produced by dissolving chromic anhydride, CrO₃, in water, or by acidifying the solution of a chromate. Its solution evolves Cl and becomes green, when it is boiled with HCl. When solution of chromic acid is neutralised by an alkali it becomes yellow, and the yellow solution yields the above reactions for chromate.

SILICATE, (SiO₃)".—Use solution of Sodium silicate, or Water-glass, Na₂SiO₃; and finely-ground sand for a solid.

Silica, or silicic anhydride, SiO_2 , occurs in nature as quartz, rock crystal, flint and sand.

645. HGl, if it is added drop by drop to solution of a silicate, which is meanwhile constantly stirred or shaken, will give a gelatinous precipitate of orthosilicic acid, H₄SiO₄, unless the solution is very dilute. Since the silicic acid may remain in solution, however, the non-appearance of a precipitate does not prove the absence of a silicate.

The solution of silicate, which is used for these reactions (1412), is made of such strength that acids do not cause a precipitate in it; but if some of the solution is boiled down to a quarter of its volume, the precipitate will usually be obtained on the addition of an acid.

646. Separation and Detection of SiO₂.—If silicate is present in small quantity, or is in dilute solution, it may be detected by acidifying the solution with HCl and evaporating the liquid to dryness in a porcelain dish. The evaporation is finished on the

water-bath (38), and the residue is heated on the water-bath as long as acid fumes are given off.

Insoluble SiO₂ is thus produced, and remains as a white powder when the dry residue is stirred and warmed with strong HCl.

During the last part of the evaporation over a flame, after the silica has separated as a gelatinous mass, spirting will occur unless the heat is lessened and the residue is stirred with a glass rod. This spirting may be also prevented by finishing the evaporation upon a water-bath (38) as soon as the liquid thickens.

The SiO_2 , which is left undissolved by HCl, is invisible as long as the liquid remains in the porcelain dish. It is rendered visible by stirring the liquid and at once pouring it out into a glass vessel. If it is free from other substances this silica will entirely disappear when it is warmed in a platinum vessel with HF-solution and strong H_2SO_4 :— $SiO_2 + 4HF = SiF_4 + 2H_2O$.

- 647. Silicates insoluble in water may be made to yield SiO_2 as is described below (a, b), the treatment depending on their behaviour with acids.
- (a) Silicates decomposable by Acid.—Many natural and artificial silicates are decomposable by evaporation with HCl, the silica being left in an insoluble condition.

The finely powdered silicate is mixed for this purpose with dilute HCl and the liquid is evaporated to dryness on the water-bath. The residue is moistened with strong HCl, then heated gently, mixed with hot water, and the liquid is boiled and filtered. White SiO₂ will be left on the filter and the filtrate will contain the metals from the silicate in solution as chlorides.

(b) Silicates not decomposable by Acid.—Many silicates belong to this class, such as mica, felspar, glass, and porcelain.

In order to remove the silica from these silicates, the finely-powdered silicate is mixed with four times its weight of fusion mixture and heated in a platinum crucible or on the foil in a powerful flame. Bubbles of carbon dioxide will be given off at first, and when these cease to escape, the temperature may be raised and the mass kept fluid for about ten minutes. It is then allowed to cool and is heated with HCl: the whole is then evaporated to dryness, the residue is moistened with strong HCl, which is then diluted with water, and the SiO_2 is separated by filtration. The metals of the silicate will now be present in the filtrate as chlorides with the alkaline chlorides.

- 648. Microcosmic-bead.—When either a silicate or SiO₂ is fused into a bead of *microcosmic s. lt*, NaAmHPO₄.4H₂O, it is not wholly dissolved, but the SiO₂ floats about in transparent particles in the melted bead, and is visible as little opaque masses when the bead is cold. The particles of SiO₂ are usually most easily seen while they are in motion in the fused bead.
- 649. Na₂CO₃ in Blowpipe-flame.—If solid SiO₂ or a silicate is heated in a fused bead of Na₂CO₃, it causes frothing: this is due to the evolution of CO₂:—Na₂CO₃ + SiO₂ = CO₂ + Na₂SiO₃.
- 650. Am₂CO₃, or AmCl, produces a gelatinous precipitate of silicic acid, H₂SiO₃, which usually forms only after a time.
- 651. Certain silicates become blue if they are moistened with cobalt nitrate solution and are then heated on charcoal in the outer blowpipe-flame.

652. Hydrogen silicate, or silicic acid, $\mathbf{H}_2\mathrm{SiO}_3$, can exist in solution in water, and the solution has a faintly acid reaction. It is ordinarily met with, however, in the hydrated state as a colourless gelatinous substance, which does not affect the colour of litmus.

• In this gelatinous form silicic acid is almost insoluble in water, and is only very slightly soluble in acids; but it dissolves readily as an alkaline

silicate, in solution of caustic alkali or of alkaline carbonate.

When the gelatinous hydrated acid is strongly heated, it yields a white insoluble powder, SiO₂, which gives the reactions in paragraphs 648, 649, and is entirely volatilised when it is warmed in a platinum vessel with HF-solution and strong H₂SO₄.

FLUORIDE, (F)'.—Use finely-powdered CaF₂.

The ordinary tests for a fluoride depend upon the liberation of hydrofluoric acid, which is then allowed to act upon a glass surface and to etch or corrode it.

653. Strong H₂SO₄, if it is warmed with a little finely-powdered fluoride in a test-tube, liberates HF. This acid etches the glass, and causes the sides of the tube to throw off the liquid when it is shaken up, as a greasy surface throws off water.

If the cooled mixture is washed out of the tube and the inside of the tube is perfectly dried, the glass will be seen to be dimmed and roughened. Gentle friction with a glass rod will often detect the roughness of the etched surface, even when this is not perceptible to the eye.

654. Etching a Watch-glass.—The preceding test is made much more delicate if it is carried out as follows.

A watch-glass is carefully heated by holding it in crucible-tongs at some distance above a flame. Its convex surface is then rubbed with a piece of paraffin-wax, and is thus covered with a melted layer. This layer is made uniform by once more heating the glass, allowing any excess of wax to drop off, and then keeping the glass moving while it cools. The coating of wax is finally hardened by immersing the glass for a short time in cold water.

Lines or characters are now traced in the wax, near the middle of the glass, by gentle pressure with the point of a penknife; the object being simply to remove the wax from these parts, and not to scratch the glass.

When the glass has been thus prepared, it is placed, with its coated surface downwards, as a cover to a platinum crucible or small leaden cup which contains a little finely-powdered fluoride and strong sulphuric acid.

Some cold water is then poured into the glass in order to keep it cool, and the bottom of the vessel is heated by a very small flame; the covered vessel is then allowed to stand for about fifteen minutes. The watch-glass is now removed, and is cleansed by holding it with crucible-tongs above the flame and quickly wiping off the melted wax.

On the clean bright surface thus obtained, the traced lines will usually be visible at once when the glass is held up to the light; but if mere traces of fluoride were present, the tracing may only become visible when the cold surface is breathed upon, and is then examined by reflected light.

655. Formation of Silicon fluoride (SiF₄).—The above methods do not serve for the detection of fluoride in the presence of silicate or of silica, since under these conditions the action of H_2SO_4 liberates SiF₄ instead of HF.

SiF₄ does not etch glass, but it gives rise to white fumes in moist air; and when it is passed into dilute AmOH, it yields a colourless flocculent precipitate of H₄SiO₄, and produces AmF in the solution.

The formation of the precipitate of silicic acid is sufficient proof of the presence of a fluoride; but after this precipitate has been filtered off, the AmF may also be detected in the filtrate. This is effected by adding CaCl₂-solution to the clear liquid, then filtering off the precipitate of CaF₂, and drying and examining it as is directed in paragraph 654.

The above method is employed for detecting fluoride in minerals which contain silica or silicate. The method is also sometimes utilised for the detection of fluoride in a substance, which has been previously intimately mixed with SiO₂ so as to render it suitable for the above test.

In order to try the reaction, a finely-powdered mixture of CaF₂ and SiO₂ may be heated with strong H₂SO₄. A small piece of marble should be dropped into the tube in which the SiF₄ is being produced: this evolves CO₂, which carries the silicon fluoride over into the AmOH-solution. The changes which occur are thus expressed by equations:—

$$2\text{CaF}_2 + \text{SiO}_2 + 2\text{H}_2\text{SO}_4 = \text{SiF}_4 + 2\text{H}_2\text{O} + 2\text{CaSO}_4.$$

 $\text{SiF}_4 + 4\text{AmOH} = \text{H}_4\text{SiO}_4 + 4\text{AmF}.$

656. CaCl₂, when it is added to solution of fluoride (*Note*), gives an almost transparent gelatinous precipitate of CaF₂, which becomes more visible when the liquid is heated or when AmOH is added: this precipitate is slightly soluble in HCl, but is almost insoluble in HĀ. [Difference from fluosilicate (470).]

Note.—Solution of NaF may be prepared for the above reaction, by heating some finely-powdered CaF₂ with melted fusion-mixture on platinum-foil for several minutes: then boiling the foil with the cool mass in water, filtering the solution, and acidifying the filtrate with HA.

657. Hydrogen fluoride, or Hydrofluoric acid, HF, at temperatures above 20 °C., is a colourless gas which fumes in the air, etches a glass surface, and dissolves freely in water. The solution of HF is acid in reaction, and differs from all other acids by decomposing and dissipating $SiO_2:-SiO_2+4HF=SiF_4+2H_2O$. It therefore also decomposes silicates and must be stored in bottles of rubber or gutta percha instead of in glass, unless the glass surface has been carefully protected by a film of wax.

Tungstate, (WO_4) .—Paragraphs 127-131.

Molybdate, (MoO_4) .—Paragraphs 236-243.

Selenite, (SeO_3) .—Paragraphs 246-250, 253, 254.

Tellurate, (TeO_4) .—Paragraphs 260, 261.

Tellurite, (TeO_3) .—Paragraphs 257-259, 261.

Titanate, (TiO_3) .—Paragraphs 333-339.

Vanadate, (VO_4) .—Paragraphs 391-397.

SECTION V

THE DETECTION AND ANALYTICAL REACTIONS OF ORGANIC SUBSTANCES.

666. Introductory Remarks.—Organic substances may be broadly defined as chemical compounds containing carbon.

When organic substances are being examined with a view to their identification, it is frequently necessary to ascertain their physical properties. Methods are therefore described in this section for determining the melting-point of a readily fusible solid, the boiling-point and density of a liquid, and the separation of liquids of different boiling-points by fractional distillation.

These descriptions are followed by an account of the methods employed for detecting the presence in organic substances of the elements carbon, hydrogen, nitrogen, sulphur, phosphorus, chlorine, bromine, and iodine.

The special analytical reactions for certain important organic substances are then described in detail. These substances include organic acid-radicles, hydrocarbons and their derivatives, alcohols and ethers, phenol and its derivatives, aldehydes and ketones, carbohydrates, glucosides, organic bases, and alkaloids.

DETERMINATION OF DENSITY, MELTING-POINT, AND BOILING-POINT.

667. The Identification of a pure Organic Substance is frequently aided by the determination of some of its constant physical properties, such as its density, and its melting-point if it is a solid or its boiling-point if it is a liquid.

These determinations may take the place of analytical tests, but they are more frequently used with the object of suggesting the analytical tests to be applied, or of confirming the results which have been yielded by the analytical reactions.

The determination of the melting-point or the boiling-point is also often of great value as a test of the purity of an organic substance. A solid substance will usually melt sharply at a particular temperature if it is pure; but if it is impure it will usually soften

before its melting-point is reached, and there may be a difference of several degrees between the softening-point and the temperature at which it becomes fluid.

Similarly if a pure liquid is distilled, the boiling-point will remain constant throughout the distillation, whereas a mixture will usually show a rise in the boiling-point as the distillation proceeds.

Accordingly these determinations will often decide whether a single substance or a mixture of substances is under examination.

The determination of the density of a solid substance involves accurate weighing on a chemical balance. Directions for the process are given in Section I. in Clowes' and Coleman's Quantitative Analysis.

The density of a liquid may be determined by accurate weighing, but the determination by means of a hydrometer (668) is much more rapid and convenient, and may be applied when a sufficient quantity of the liquid is available.

When a solid organic substance possesses a melting-point which can be observed on a mercurial thermometer, the process described in paragraphs 669 and 670 may be employed, and will be found to be rapid and simple in execution.

The determination of the boiling-point of a simple liquid, if the temperature falls within the range of a mercurial thermometer, is described in paragraph 671.

Fig. 58.

The constituent liquids of a mixture may often be separated from one another by fractional distillation (672), and the separated liquids may then be identified by their boiling-points or by other methods.

DETERMINATION OF THE DENSITY OF A LIQUID.

668. The Hydrometer serves to determine rapidly the density of a liquid, since it enables the process of weighing to be dispensed with. The apparatus consists of a glass or metal float (Fig. 58), which is weighted below to such an extent as to cause it to assume a vertical position when it is floating in the liquid. The stem is so graduated that the numbered mark, which is level with the surface of the liquid when the hydrometer is floating freely, shows the density of the liquid.

The temperature of the liquid must be adjusted to the temperature at which the hydrometer has been graduated, which is usually 15.5° C.; and all densities The Hydro of liquids which are given in this book correspond to METER.

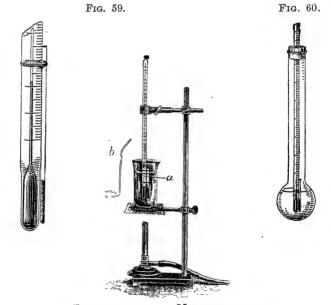


that temperature, and are referred to the density of distilled water at 4° C. as unit, unless it is otherwise stated.

DETERMINATION OF THE MELTING-POINT OF A SOLID.

Two methods, which are commonly in use for determining the melting-point of a substance, are described below.

669. First Method.—The solid substance is placed in a thinwalled glass tube, which is about 1 mm. in diameter and is sealed at one end. Such a tube is readily made by drawing out a thinwalled narrow test-tube in the Bunsen-flame. The long tube thus



DETERMINATION OF MELTING-POINT.

produced is cut into tubes 5 cm. in length, each of which is then closed at one end by fusion in the flame.

The open end of one of these tubes is dipped into the finely powdered substance, and the tube is placed mouth upwards and gently tapped to cause the powder to fall to the closed end. These operations are repeated until about 1 cm. of the tube is filled with the powder.

The tube is then attached closely to the stem of a delicate thermometer with the powder alongside the bulb, by binding it with fine platinum wire (Fig. 52, a). If the thermometer is to be dipped into water only, a little rubber ring, cut off some small rubber tubing,

may be slipped over the larger end of the tube and the thermometer stem, instead of binding it with platinum wire.

The thermometer and tube are then placed in a beaker containing a liquid which has a higher boiling-point than the melting-point of the solid. Water, strong-sulphuric acid, or melted paraffinwax may be used in the beaker, according to the temperature which is required.

The beaker is then heated until the solid melts, and the temperature of liquefaction is accurately noted by the thermometer.

During the process of heating, the contents of the beaker should be constantly mixed by means of a stirrer, which is made of bent glass rod or stout wire in the shape shown at b in Fig. 59. This continuous stirring is necessary, in order to secure a uniform temperature throughout the mass of the liquid during the determination.

The process should be repeated several times with fresh portions of the substance, and the mean temperature, which has been obtained from these experiments, should be taken as the melting point of the solid. The temperature of solidification may also be noted and utilised in most cases as a check.

If the mercury-thread of the thermometer extends above the surface of the liquid, the correction which is described in the *Note* in paragraph 671 must be applied to the reading.

In the case of certain fats, and of some other substances, it is difficult to see the exact point of liquefaction. The observation is facilitated by using a tube which is open at both ends, but it is essential in this case to select a liquid for the beaker, which does not act chemically upon the substance.

If there is any difficulty in introducing the substance into this open tube, the fine end of the tube may be inserted into the melted substance, when a small quantity of the liquid will at once enter the tube by capillary attraction. This is allowed to cool and solidify, and the tube is then attached to the thermometer and immersed in the water as before.

At the moment of liquefaction, the substance will be forced up this open tube by the pressure of the liquid in the bath, and the temperature of the thermometer is noted when this movement is seen to occur.

It should be borne in mind that some substances, more especially the animal fats, show a lower melting-point when the process of fusion is repeated immediately after solidification. In such cases the normal melting-point is regained after the substance has remained in the solid condition for a sufficient interval of time.

EXPERIMENT.—The melting-point of naphthalene may be determined using a beaker which contains water for the bath. The melting-point should be approximately 80° C.

670. Second Method.—A modified form of vessel for determining melting-points is shown in Fig. 60. It is made by blowing a bulb at the end of a piece of wide glass-tube, which should be about 15 cm. in length, with a diameter of about 3 cm. and the bulb should be about 5 cm. in diameter. This bulb is nearly filled with strong sulphuric acid.

The bulb of the thermometer is then wetted with the acid by dipping it beneath the surface (Fig. 60), and the tube containing the substance is placed alongside and in close contact with the wet thermometer bulb as is shown in Fig. 59, α ; the tube will adhere to the wetted bulb.

The thermometer bulb with the adhering tube is then immersed in the acid, the upper part of the tube remaining above the surface of the liquid; the small tube will continue to adhere to the thermometer owing to the surface-tension of the acid film. The stem of the thermometer is held in this position by means of a cork inserted in the mouth of the large tube, the cork having a groove cut in its edge to permit the air to escape as it is expanded by heat.

When the apparatus has been properly adjusted, heat is gradually applied to the bulb, and the melting-point is observed. The process is repeated, and the results obtained are dealt with as has been already described in the first method (669).

The discoloration of the acid, which may be caused by its continued use, may be removed by dropping in a crystal of potassium nitrate.

EXPERIMENT.—Determine the melting-point of sulphur or of benzoic acid by the above method.

DETERMINATION OF THE BOILING-POINT OF A LIQUID.

671. The Boiling-point of a simple Liquid may be determined in the following way.

The liquid is placed in a flask such as is shown in Fig. 61, the neck of which is closed by a perforated cork through which a thermometer passes, the bulb of the thermometer being close to the surface of the liquid. The liquid is then heated to boiling, and the bulb and part of the stem of the thermometer are thus surrounded by the vapour of the boiling liquid.

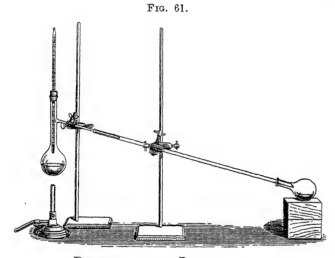
The side exit-tube in the neck of the flask should be connected with a cold-water condenser, if the liquid has a low boiling-point; but a plain glass tube, about thirty inches in length and cooled only by the atmosphere, will suffice for less volatile liquids (Fig. 61).

As soon as the reading of the thermometer remains constant, the temperature is registered as the boiling-point of the liquid.

Note.—This reading will, however, require to be corrected if the thread of mercury in the thermometer extends above the cork. The following formula gives the correction to be added:—

$$N(T - t) \times 0.000156.$$

Here N is the number of degrees on the thermometer-stem which are not heated by the vapour, T is the temperature indicated by the thermometer, t is the temperature taken by placing the bulb of a second thermometer midway between the top of the cork and of the mercury column during the



DETERMINATION OF BOILING-POINT.

process, this second bulb being screened from the direct heat of the Bunsen-flame, and 0 000156 is the apparent coefficient of expansion of mercury.

It should be remembered that considerable differences of atmospheric pressure will cause appreciable variations in the boiling-point. All the boiling-points which are given in this book correspond to the normal atmospheric pressure of 760 mm. of mercury.

EXPERIMENT.—The boiling-point of aniline may be taken by this method. It should be approximately 78° C., and a cold-water condenser will therefore not be required.

SEPARATION OF LIQUIDS BY FRACTIONAL DISTILLATION.

672. Fractional Distillation may be employed to separate from their mixture two or more muscible liquids, which differ in their boiling-points by at least 20 degrees.

If a mixture of two such liquids is placed in a distillation flask (Fig. 61), fitted with a thermometer and a cold-water condenser,

and the mixture is gradually heated, the following results will be observed.

The liquid will begin to boil at about the boiling-point of the more volatile liquid, but as the heating is continued the boiling-point will gradually rise. After a time the boiling-point will rise rapidly until it approaches the boiling-point of the less volatile liquid, when it will again rise slowly until the whole of the liquid has distilled.

If the volume of the distillate is measured from time to time, it will be found that the two largest volumes have been obtained while the thermometer indicates approximately the boiling-points of the two liquid constituents of the mixture; and that a partial separation of the constituents has therefore been effected, if the separate fractions of the distillate which come over at fairly constant temperatures have been received in separate vessels.

If each of these two main "fractions" is now separately distilled or "refractionated," it may be separated in a similar manner into fractions; and a fairly pure sample of each of the constituent liquids may be obtained by again collecting separately those fractions of the distillate which pass over while the thermometer registers the boiling-point of the particular constituent.

By a further "refractionation," or repetition of this fractional distillation, a pure specimen of each of the original liquids may ultimately be obtained.

The following example will illustrate this method of separation by fractional distillation.

EXPERIMENT.—Make a mixture of equal volumes of pure alcohol and water. Pour 150 c.c. of this mixture into the distillation-flask fitted with a thermometer and a cold-water condenser: heat the liquid gently with a naked flame, and receive the fractions which pass over between 78°–85°, 85°–92°, 92°–98°, and 98°–100° separately in four clean dry flasks.

When the volumes of these fractions A, B, C, and D, are measured, fraction A will be about one-third of the total volume, and may be proved by its smell and by its ready inflammability to contain the greater part of the alcohol.

The fractions B and C will be smaller in volume and will consist of a mixture of alcohol and water, the proportion of alcohol being so small that the mixture is not inflammable.

Fraction D will be found by its tastelessness and freedom from smell to consist almost entirely of water.

A further separation may be effected by repeating the above procedure with the fractions already obtained, until eventually the fractions with the lowest boiling-point will consist of almost pure alcohol, and those with the highest boiling-point will consist of almost pure water.

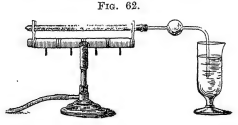
It will be obvious that a separation of the liquids in the mixture has thus been effected, and that special methods may now be applied for the identification of each of the liquids.

This method of procedure is often applicable to liquid mixtures, the composition of which it would be difficult or impossible to ascertain until it becomes possible to deal with each constituent of the mixture in the separate condition.

DETECTION OF THE ELEMENTARY CONSTITUENTS OF ORGANIC SUBSTANCES.

DETECTION OF CARBON.

- 673. Ignition of Substance.—The presence of carbon in an organic substance is frequently indicated, by the substance becoming charred or blackened, when it is heated alone or with strong $\rm H_2SO_4$. The carbon may also be evolved as $\rm CO_2$ when the substance is heated either alone or with strong $\rm H_2SO_4$, and the $\rm CO_2$ thus evolved may be detected by means of lime-water (474).
- 674. Ignition with Copper oxide.—A more certain and general method of detecting carbon consists in mixing the organic substance intimately with finely-powdered copper oxide (CuO), or



DETECTION OF CARBON AND HYDROGEN IN AN ORGANIC SUBSTANCE.

with some other solid oxidising compound, and heating the mixture strongly. The carbon is then wholly evolved as CO₂, and this may be detected by passing it into lime-water (474).

If perfectly dried materials are used (Note, 676) the process serves also for the detection of hydrogen as a constituent of the organic substance (676).

The method may be tried by mixing a little sugar with four or five times its measure of finely-powdered CuO. This mixture is then placed in the closed end of a hard glass test-tube, and the remainder of the tube is nearly filled with granular CuO. The open end of the test-tube is then fitted with a bent glass delivery-tube, and the tube is heated in a Bunsen-flame, care being taken that the granular oxide is heated to redness before the mixture is heated.

The experiment may be made in a longer tube, which is heated by the Ramsay-burner (Fig. 62), the heat being first applied to the granulated oxide, and when this is at a red heat the mixture being gradually heated to the same temperature. The delivery-tube dips into a vessel containing lime-water, which serves to indicate the evolution of CO_2 .

Note 1.—In the case of a volatile liquid the apparatus must be modified so as to cause the vapour of the liquid to pass slowly over the red-hot copper oxide, the products of combustion passing on into the lime-water.

Note 2.—It must be remembered that cyanates, many carbonates, and all bicarbonates, evolve CO_2 when they are heated. Carbonates may be distinguished from other organic bodies by evolving CO_2 when they are acted upon by cold dilute acid (474).

DETECTION OF HYDROGEN.

675. Ignition of Substance.—Many organic substances, which contain both hydrogen and oxygen, evolve their hydrogen more or less completely in the form of steam when they are heated. If the dry substance is heated in a test-tube, and steam is evolved which condenses on the cooler part of the tube in the form of water drops, the substance is proved to contain hydrogen and oxygen.

676. Ignition with Copper oxide.—The presence of hydrogen in a dry organic substance may, however, be detected with greater certainty at the same time as the carbon, by the method of oxidation with dry CuO (see *Note*) at high temperature (674).

The hydrogen is completely converted by this oxidation process into steam, which condenses in the form of water drops in the cooler part of the test-tube, or in the delivery-tube. The bulb in the delivery-tube (Fig. 62, p. 181) is of advantage, since it reduces the velocity of the current of escaping gas and vapour and also exposes it more fully to the cooling action of the air, and therefore promotes the condensation of the steam.

The presence of hydrogen in sugar will have been already proved, if perfectly dry materials were used, by the production of the water,

which has condensed in the apparatus during the test for carbon (674). The test is rendered more delicate by placing some white anhydrous CuSO₄ in the bulb, since the white powder becomes blue in the presence of even a small amount of water.

Note.—Any moisture which is present in the substance or in the copper oxide will be evolved as steam, and will be condensed as is described above. Both the organic substance and the CuO must therefore have been perfectly dried, in order to prevent a wrong inference from being drawn as to the presence of hydrogen in organic combination in the substance.

A solid organic substance may usually be freed from moisture by heating it in the steam-oven (48) for about an hour, and the copper oxide should be dried by igniting it to redness for an hour and cooling it in a desiccator.

DETECTION OF NITROGEN.

- 677. Ignition of Substance.—Nitrogen is evolved from many organic substances in the form of ammonia, or of an alkaline organic derivative of ammonia, when the substance is heated. The ammonia, or other alkaline substance, may usually be readily detected by its pungent smell or by its alkaline action upon moist turmeric-paper (*Note*, 678).
- 678. Ignition with Soda-lime.—A general method of detecting the presence of nitrogen consists in mixing the organic substance intimately with powdered soda-lime, and then heating the mixture strongly in a hard glass test-tube. Either ammonia or a basic substance containing nitrogen, such as an amine, will be evolved, and may be detected by turning moist turmeric-paper brown.

In trying the reaction urea or albumen may be employed.

This test is applicable to most organic substances which confidences.

This test is applicable to most organic substances which contain nitrogen, with the exception of nitro-, nitroso-, azo-, and diazobodies.

Note.—It must be remembered that many ammonium compounds evolve NH_3 when they are heated, and that they all evolve NH_3 when they are heated with soda-lime. When ammonium compounds, however, are mixed with solution of KOH, they evolve NH_3 at ordinary temperatures or when they are gently heated; and they may be distinguished in this way from most organic compounds containing nitrogen, since these require to be heated much more strongly with solution of KOH, and in some cases with solid KOH or with soda-lime, before they evolve NH_3 .

679. Ignition with Sodium and Production of Prussian. Blue.—Another general method for the detection of organic nitrogen consists in converting the nitrogen of the substance into a soluble cyanide, and the cyanide into "Prussian blue."

The procedure is described in the following paragraph, and either urea or albumen may be used as the nitrogenous organic substance.

Heat the nitrogenous substance strongly with a small piece of

sodium or potassium in a little hard-glass ignition-tube (see *Note*) When the substance has been charred, break up the tube in some water in a mortar with the pestle, transfer the liquid with the substance to a boiling-tube, warm gently and filter. Then add a few drops of solutions of FeCl₃ and FeSO₄: and unless the liquid is already strongly alkaline to test-paper, add KOH in excess. Then boil the liquid, and add HCl in excess.

The formation of a dark blue precipitate, or of a blue or dark green solution, indicates the formation of Prussian blue; and the nitrogen, which is a necessary constituent of this Prussian blue, must have been derived from the organic substance.

This reaction is more delicate than the preceding one (678), and it serves for the detection of nitrogen in all organic substances with the exception of the diazo-compounds.

Note.—If sulphur is present, the organic substance must be mixed with iron filings, or better with "reduced iron," before it is heated with sodium.

The absence of cyanogen-compounds in the organic substance must be ascertained (692 and 694) before applying this test for nitrogen.

DETECTION OF SULPHUR.

680. The Presence of Sulphur in an organic substance may be detected by converting the sulphur either into Na₂S (681), or into Na₂SO₄ (682).

Albumen may be used in trying the following tests.

681. Ignition with Sodium, and formation of Sulphide.— Ignite the organic substance with a small piece of sodium: then extract the residue with warm water, filter the solution, and add to the filtrate, made alkaline if necessary with NaOH, some solution of sodium nitroprusside: the presence of sulphur will be indicated by a violet coloration (483).

Lead acetate-solution may be added instead of nitroprusside, and will produce black PbS (482); but this is a less delicate test than the nitroprusside reaction.

682. Ignition with NaNO₃, and formation of Sulphate.— Mix the solid organic substance in a porcelain crucible with four times its measure of dry Na₂CO₃ and its own measure of NaNO₃, and heat the mixture until the alkali salts are fused. Dissolve the cool product in dilute HCl, and add BaCl₂: a white precipitate of BaSO₄ (464) shows the presence of sulphate, which has been formed from sulphur in the organic substance.

DETECTION OF PHOSPHORUS.

- 683. The Presence of Phosphorus in an organic substance may be detected by converting the phosphorus either into PH₃ (684), or into phosphate (685).
- 684. Heating with Magnesium.—Char the substance by heat, then mix it with magnesium-powder and heat the mixture strongly. Now moisten the cold residue with water: a garlic-like smell of PH₃ will be observed, showing that magnesium phosphide has been produced during the ignition, and has then been decomposed by the water. This proves the presence of phosphorus in the original substance.
- 685. Ignition with NaNO₃, and formation of Phosphate.—Heat the solid substance with fused Na₂CO₃ and NaNO₃, as has been described for the detection of sulphur (682). Then dissolve the residue in dilute HNO₃, and add a few drops of this liquid to some AmHMoO₄-solution and warm gently: the formation of a yellow precipitate (571) proves that phosphate has been formed, and that phosphorus was present in the original substance.

DETECTION OF CHLORINE, BROMINE, AND IODINE.

686. The Presence of the Halogen Elements in an organic substance may be detected either by the flame-coloration of their copper salts (687), or by converting the halogens into sodium salts and then applying the ordinary tests to the solution of these salts (688).

Chloral may be used for trying these methods of detection of organic chlorine.

687. Copper Flame-coloration.—Fix a piece of CuO in a loop of platinum wire (*Note*, below), and heat it strongly in the outer Bunsen-flame until it gives no colour to the flame. Then dip the CuO into the organic substance and ignite it again, first in the inner and then in the outer Bunsen-flame: a green flame-coloration with a blue centre will appear, and will indicate the presence of chlorine in the organic substance.

A similar result is obtained if the organic substance contains either bromine or iodine. This test, therefore, only serves to indicate the presence of a halogen, and does not distinguish between the different halogen elements.

Note.—The expensive platinum wire may be replaced by a piece of fine copper wire: this must be superficially oxidised by heating it in the outer flame, and is then dipped into the organic substance.

688. Ignition with Soda-lime or with Sodium.—Mix a little of the organic substance with soda-lime free from chloride, place the mixture in a hard glass test-tube, and add a layer of soda-lime. Then heat the soda-lime to redness (Fig. 62 p. 181), and gradually extend the heating to the mixture until the whole is red hot. Now allow the whole to cool, and dissolve the cold soda-lime in dilute nitric acid. Test this clear solution for chloride, bromide and iodide in the ordinary way (539-556).

The substance may be ignited with a small piece of sodium, instead of with soda-lime, and the clear solution obtained by extracting the cold residue with water is then examined for the halogens in the ordinary way, 539-556. It must be remembered that the process of ignition with sodium will produce a cyanide if nitrogen was present in the substance, and that this might be mistaken for a halide (679, 691, et seq.).

ANALYTICAL REACTIONS OF ORGANIC SUBSTANCES.

GROUP I.—ORGANIC ACID-RADICLES.

1. CYANOGEN ACID-RADICLES.

This Group of acid-radicles includes cyanides, cyanates, thio-cyanates, ferrocyanides and ferricyanides.

CYANIDE, (CN', or Cy').—Use freshly-made solution of KCN.

The alkaline cyanides smell faintly of bitter almonds, the smell being due to the liberation of HCN by the action of the ${\rm CO_2}$ and ${\rm H_2O}$ in the air. Hydrocyanic acid is intensely poisonous.

- 689. Dilute acids liberate HCN with an odour of bitter almonds; great care should be taken not to inhale this acid vapour to any extent, it should be only cautiously wafted to the nose with the hand. Certain complex cyanides, such as K₄FeCy₆ and K₃FeCy₆, also evolve HCN when they are treated with dilute acid (705, 712).
- 690. Strong H₂SO₄, when it is warmed with any cyanide, whether it is simple or complex, causes decomposition with evolution of CO, which burns with a blue flame.

691. AgNO3: a white precipitate of AgCN.

This precipitate is most easily obtained by pouring a drop of the KCN-solution into the AgNO₃-solution.

Show with three separate portions of the precipitate in the liquid, that the precipitate is soluble in K(N-solution when it is added in excess, and in AmOH, but is insoluble in dilute $HN()_3$.

Filter off some of this precipitate, and wash it with hot water, until it is proved to be free from AgNO₃ by HCl causing no milkiness with the last few drops of the washing water. Then dry it, scrape it from the filter into a porcelain crucible, and heat it strongly; metallic silver will remain. If the cool residue is warmed with a little dilute HNO₃ it will dissolve, giving off reddish brown gas; and when a drop of HCl is added to this solution, a curdy precipitate of AgCl will be produced.

This method of examining the precipitate which has been produced by AgNO₃ distinguishes AgCN from AgCl, AgBr, and AgI; these are also precipitated by AgNO₃ and are insoluble in HNO₃, but when they are dried and strongly heated, they fuse without undergoing decomposition. Hence, when HNO₃ is warmed with the cool mass after fusion, it cannot dissolve out Ag;

and when HCl is added to the HNO₃, it either gives no precipitate, or it causes only a slight milkiness, which is due to the precipitate not having been perfectly freed from soluble silver salt.

This process of proving the silver precipitate to consist of AgCN is somewhat tedious and need seldom be used, since cyanide is readily detected and

distinguished by the following reactions.

692. "Prussian Blue" test.—FeSO₄-solution and FeCl₃-solution are added to solution of cyanide, and the liquid is then made distinctly alkaline with KOH, and boiled; it is then cooled and acidified with 'HCl: "Prussian blue," Fe₄(FeCy₆)₃, will be formed, and will appear as a deep blue precipitate: or if the quantity of cyanide originally present was small, a bluish-green solution of the Prussian blue will be obtained.

If no cyanide is present, the addition of HCl will produce an almost colourless and perfectly clear liquid.

693. Formation of Thiocyanate, or Sulphocyanide, AmCNS.—If dilute H₂SO₄ is added to a cyanide in a test-tube fitted as is shown in Fig. 56, 474, and then a small piece of marble is dropped in and the liquid is boiled, HCN will be carried over by the CO₂ gas; and if the gas is allowed to bubble through some Am₂S-solution, AmCNS will be formed.

If the Am₂S-solution is now boiled in a porcelain dish, and acidified when cold with HCl in order to remove unchanged Am₂S, the addition of several drops of FeCl₃ will produce a red colour (703). The colour will be easily seen against the interior of the white dish. This result proves that HCN has passed into the Am₂S, and therefore that cyanide was present in the original substance.

A little of this red liquid should be added to some HgCl₂-solution. If the red colour is destroyed, the presence of a thiocyanate is confirmed.

When HCl is added to the Am₂S in the above reaction, white S will separate. If the yellow liquid has been previously boiled in a porcelain dish until it becomes colourless, filling in distilled water to prevent evaporation to dryness, the separation of S on the addition of an acid is prevented. But this separation of S is frequently advantageous, since its whiteness helps to show up even a faint reddish tint in the liquid.

The marble is added in order to cause evolution of CO₂ gas: the escape of the gas lessens the risk of the Am₂S being sucked back during the reaction: it also helps to carry over the HCN into the Am₂S-solution, and to prevent it from remaining behind in the test-tube.

694. The formation of Thiocyanate can often be more simply secured, by boiling the solution of the cyanide in a porcelain dish with yellow Am₂S. The Am₂S must be added until the yellow colour of the mixture remains after the liquid has been boiled for a short time. If a dark-coloured precipitate is formed, the yellow colour of the liquid can only be seen after the precipitate has been

removed from a portion of the liquid by settling or by filtration. The whole of the liquid is then filtered, if necessary, and is acidified with HCl in the porcelain dish, and FeCl₃ is added in small quantity: a blood-red colour will be produced, which will not be removed when the liquid is heated or is mixed with a little dilute HCl. The colour will be at once destroyed, however, when a few drops of the liquid are added to some HgCl₂-solution.

Note.—The methods described in paragraphs 692, 693, 694 serve as exceedingly delicate tests for cyanide. The methods in 692 and 693 are often less readily performed than that in 694, but they exceed it in delicacy. The method in 694 is only replaced by that in 693 in cases where the filtrate is dark coloured after the substance has been boiled with an excess of Am₂S, and the red coloration would therefore be invisible.

The results of the reactions in paragraphs 692, 693 and 694 are also given by many complex cyanides.

695. HgCy₂ and AgCy do not show the Reactions for Cyanide by methods 691, 692, 693, but when they are boiled with Am₂S, according to method 694, AmCNS will remain in the filtrate from the black precipitate of sulphide, and will give the red coloration with FeCl₃.

When solid HgCy₂ is heated in a small tube closed at one end, it gives off cyanogen gas, which burns with a peach-blossom coloured flame if it is kindled. This gas is very poisonous when it is inhaled.

696. Hydrogen cyanide, Hydrocyanic acid or Prussic acid, HCN, is a colourless, volatile, inflammable liquid which boils at 26° C. It readily dissolves in water, and the vapour emitted from the acid or its solution smells like bitter almonds. The acid is highly poisonous when it is inhaled as vapour or swallowed as liquid. Hydrocyanic acid is detected by (691-694).

CYANATE, (CNO)'.—Use a freshly-made cold solution of KCNO.

697. The cyanates of the alkalis and alkaline earths are stable in the dry state, but they change in moist air or in solution to a mixture of bicarbonate and ammonia.

698. Dilute Sulphuric acid sets free cyanic acid, which gives an irritating and pungent odour. The acid decomposes into an ammonium salt and carbon dioxide:—HCNO + H₂O = CO₂ + NH₃; hence CO₂ mixed with a small amount of pungent HCNO is given off with effervescence, and the solution contains Am_2SO_4 , which will evolve ammonia when it is warmed with KOH.

699. AgNO₃: white curdy precipitate, AgCNO, soluble in AmOH and in HNO₃. (Difference from cyanide.)

700. Cobalt acetate is coloured blue by a solution of the cyanate, owing to the formation of blue potassium cobaltocyanate, K_2 Co (CNO)₄.

THIOCYANATE, OR SULPHOCYANIDE, (CNS)'.—Use solution of KCNS.

701. Moderately strong H2SO4 decomposes thiocyanates with

evolution of COS, which burns with a blue flame :—KCNS + $\rm H_2SO_4$ + $\rm H_2O$ = COS + KHSO₄ + NH₄HSO₄.

702. AgNO₃: white curdy precipitate, AgCNS, soluble in AmOH, insoluble in HNO₃.

703. FeCl₃: blood-red coloration, but no precipitate: the colour is weakened but not destroyed by HCl: it disappears entirely when the liquid is dropped into HgCl_2 -solution. [Difference from similar colorations produced by other acid-radicles.]

704. CuSO₄, added drop by drop, gives first an emerald green liquid and then a black precipitate of Cu(CNS)₂.

If sulphurous acid is then added, white CuCNS is produced from the black Cu(CNS)₂. If H₂SO₃ is added before the CuSO₄ to the sulphocyanide solution, the white precipitate is at once produced.

FERROCYANIDE, (FeCy₆)^{IV}.—Use solution of K₄FeCy₆.3H₂O.

705. Dilute H₂SO₄ produces no change at ordinary temperatures; but when it is heated to boiling, HCN is evolved:—

 $2K_4FeCy_6 + 3H_2SO_4 = K_2Fe.FeCy_6 + 3K_2SO_4 + 6HCN.$

706. Strong H₂SO₄ decomposes ferrocyanides completely with evolution of CO:—

 $K_4 \text{FeCy}_6 + 6H_2 \text{SO}_4 + 6H_2 \text{O} = 6\text{CO} + 2K_2 \text{SO}_4 + \text{FeSO}_4 + 3\text{Am}_2 \text{SO}_4.$

. 707. Strong HCl: white precipitate of H₄FeCy₈, which gradually changes to blue.

708. FeCl₃: dark blue precipitate of "Prussian blue," Fe₄(FeCy₈)₃, insoluble in HCl, soluble in H₂C₂O₄ to a dark blue liquid, and changed by KOH into brown Fe(OH)₃.

The solubility in $\rm H_2C_2O_4$, and the insolubility in HCl, may be shown by pouring off portions of the liquid containing the precipitate, heating them with HCl and with $\rm H_2C_2O_4$ respectively, then filtering and noting that the filtrate is rendered blue by the solution of the precipitate.

709. FeSO₄: light blue precipitate, K₂Fe(FeCy₆); which becomes darker in colour by oxidation on exposure to the air, or on the addition of Cl- or Br-water, or when it is warmed with HNO₃, owing to the formation of Prussian blue (708).

710. CuSO₄: a chocolate-coloured precipitate, Cu₂FeCy₆, insoluble in HA. This is very characteristic.

711. AgNO₃: white precipitate, Ag₄FeCy₆, insoluble in HNO₃

and in AmOH. When the precipitate is heated with HNO₃, it changes to orange-red Ag₃FeCy₆ which is soluble in AmOH.

FERRICYANIDE, (FeCy₆)"".—Use fresh K₃FeCy₆ solution.

A ferricyanide may be prepared by oxidising a ferrocyanide by means of Cl, Br, ${\rm H_2O_2}$ or other suitable oxidant.

- 712. Dilute and strong H₂SO₄ behave with ferricyanides in the same manner as with ferrocyanides (705, 706).
- 713. FeCl₃: dark green or brown coloration; no precipitate is formed, the liquid becoming transparent on dilution.
- 714: FeSO₄: dark blue precipitate of "Turnbull's Blue," Fe"₃(FeCy₆)₂, insoluble in acids: KOH destroys the blue colour.
 - 715. CuSO₄: yellowish precipitate.
- 716. AgNO₃: orange-coloured precipitate, Ag₃FeCy₆, insoluble in HNO₃, soluble in AmOH: a white residue undissolved by AmOH proves that some K₄FeCy₆ was originally present (711).
- 717. Detection of Ferro- and Ferricyanide.—Ferro- and ferricyanide can be found and distinguished by pars. 708–710 and 713–715, or by the precipitate given by AgNO₃ in the solution acidified with HNO₃. This precipitate will have a more or less decided orange-red colour if a ferricyanide is present.

If the precipitate is separated from most of the liquid by filtration or decantation and is gently heated with AmOH, white Ag_4FeCy_6 will be left if a ferrocyanide was present, and when the filtered ammoniacal liquid is acidified with HNO_3 orange-red Ag_3FeCy_6 will be reprecipitated.

Another method for detecting cyanide, ferrocyanide, ferricyanide and thiocyanate in the presence of one another is given in paragraph 1284.

725. Introductory Remarks on other Acid-radicles.—The other acid-radicles whose reactions are described in this Section are all radicles of carboxylic acids, with the exception of urates (807).

Pyrogallic acid, picric acid, and carbolic acid are more correctly sermed pyrogallol, trinitrophenol and phenol respectively: and their seactions are described under phenol derivatives (848-871), as shey are not true acids.

Carboxylic acids are characterised by containing the monovalent earboxyl group COOH, in which the hydrogen atom is replaceable by a metal. These acids are usually weak acids, and are only elightly dissociated by solution.

In aqueous solution the organic acid R:COOH is dissociated into

the ions H and R COO': thus acetic acid, CH_3 .COOH, is separated into the ions H and CH_3COO' , and succinic acid, $C_2H_4(COOH)_2$, into the ions 2H and $C_2H_4(COO)_2''$. It is evident therefore that the basicity of the acid corresponds to the number of COOH-groups which are present in its molecule.

The organic acids themselves are often available for trying the reactions: the corresponding salts of the alkali-metals are usually obtainable, and less frequently the salts of organic bases or of heavy metals. The tests may be applied directly to the salts of the alkali-metals: but the free acids must be neutralised with sodium hydroxide, since a neutral solution is used when possible for trying the reactions.

In the following statement the general character and properties. of each acid are first described, and then the most important reactions of its salts are given.

2. ALIPHATIC ACID-RADICLES.

The acid-radicles of this class, whose reactions are described are formate, acetate, oxalate, succinate, malate, tartrate and citrate,

FORMATE, (CHO₂)'.—Use Sodium formate, NaCHO₂. Formic acid and formates much resemble acetic acid and acetates.

726. Formic acid, H.COOH, is a colourless fuming liquid, with a pungent smell; it boils at 100.6° C., has a density of 1.2256, and is readily miscible with water, with alcohol, and with ether. The acid and its salts are powerful reducing reagents.

All the formates, with the exception of the formates of Pb and Hg, are readily soluble in water.

- 727. Ignition.—A solid alkaline formate fuses and chars and emits CO and CO₂, when it is strongly heated. If it is heated with soda-lime hydrogen is evolved.
- 728. H_2SO_4 : when a formate is heated with strong H_2SO_4 , it evolves CO gas, which burns with a blue flame. No charring or blackening occurs, and no CO₂ is evolved, as in the case of an oxalate (739).

If a formate is distilled with dilute H₂SO₄, a distillate is obtained which emits the pungent smell of formic acid.

When the red liquid containing excess of formate is boiled, the Fe is completely precipitated as basic ferric formate, and the liquid becomes colourless. This reaction is precisely similar to that for acetate (736).

730. Silver mirror.—When solution of a formate is gently heated with ammoniacal solution of AgNO₃ (Note, 759), it gives no silver mirror. But free formic acid, or solution of a formate which has been acidified with HĀ, gives a silver mirror when it is heated with AgNO₃-solution.

√731. HgCl₂, when it is mixed with solution of a formate and heated to about 70° C., yields a white precipitate of Hg₂Cl₂, if HCl and alkaline chlorides are absent.

/ACETATE, $(C_2H_3O_2)'$ or $(\bar{\Lambda})'$.—Use Sodium acetate, Na $\bar{\Lambda}$.3 H_2O .

732. Acetic acid, CH₃.COOH, is known as glacial acetic acid. At temperatures below 16·7° C. it is a colourless crystalline substance, which melts at 16·7°, and boils unchanged at 118° C., giving off a very pungent inflammable vapour. This acid does not redden blue litmus test-paper until it is diluted. It is miscible with water, with alcohol, and with ether in all proportions.

The acetates, with the exception of acetates of Hg and Ag and a few basic acetates, are soluble in water.

733. Ignition.—Many solid acetates, when they are strongly heated, blacken and evolve the vapour of acetone; this vapour possesses a peculiar smell, and burns with a bright flame when it is kindled.

√734. Strong H₂SO₄, when it is heated with an acetate, evolves vapour of HĀ. This vapour, if it is smelt in an undiluted state, is very pungent; in mixture with air it smells like vinegar.

735. Formation of Ethyl acetate.—If an acetate is first mixed with a little alcohol, and then with strong H_2SO_4 , a fragrant smell of ethyl acetate, $C_2H_5\bar{A}$, will be emitted when the mixture is warmed. The smell is most pronounced when the liquid is allowed to cool for a short time, and is then shaken up.

736. FeCl₃, if it is added in small quantity to solution of an acetate, causes a deep red coloration.

The red colour of the solution is changed to yellow by the addition of a few drops of dilute HCl; it is distinguished in this way from the coloration which has been caused by thiocyanate (703).

If the red liquid contains an excess of the acetate and is boiled, the Fe is completely precipitated as light brown basic acetate, and the liquid becomes colourless.

737. Cacodyl reaction.—This reaction must be carried out with due caution, since the vapour of cacodyl is very poisonous.

When a solid acetate is heated with a minute quantity of As₄O₆ the disgusting smell of the vapour of cacodyl, As(CH₃)₂, is evolved.

OXALATE, (C₂O₄)".—Use Ammonium oxalate, Am₂C₂O₄.2H₂O.

738. Oxalic acid, $(COOH)_2.2H_2O$, crystallises in colourless rhombic crystals, which are soluble in water and in alcohol, and slightly soluble in ether: the aqueous solution is poisonous. The acid melts at 98° C., and at 150° C. it partly sublimes unaltered evolving vapour which causes coughing, and is partly decomposed:— $H_2C_2O_4 = H.COOH + CO_2$. It yields the reactions in paragraphs 739-743.

Most oxalates are insoluble in water.

739. Strong H_2SO_4 , if it is heated with a solid oxalate, causes an effervescence, which is due to the evolution of a mixture of CO and CO_2 :—

 $Am_2C_2O_4 + 3H_2SO_4 = CO + CO_2 + 2AmHSO_4 + H_2SO_4 \cdot H_2O$.

The CO may be detected by burning with a blue flame when a light is applied, and the CO₂ is found by rendering lime-water milky (474, 1, 3, 4).

No blackening or charring is caused by strong H₂SO₄. Oxalates differ in this respect from most other organic acids and their salts.

- 740. Solution of CaCl₂, of CaSO₄, or of CaH₂O₂, when it is added to solution of an oxalate, gives a white precipitate of CaC₂O₄, which is insoluble in AmOH and in HĀ, but soluble in HCl and in HNO₃. Keep this precipitate.
- 741. Conversion into Carbonate by Ignition.—Filter off the above precipitate of CaC₂O₄ (740), and pour upon part of it some dilute HCl; it will not effervesce.

Dry the remainder of the precipitate, and heat it for a moment to dull redness on platinum foil; it will be converted without blackening into CaCO₃, and this will effervesce when dilute HCl is poured upon it.

This is a general test for oxalate of K, Na, Ba, Sr, or Ca; since each of these oxalates, when it is ignited, leaves a carbonate which effervesces with an acid. Many other organic salts of these metals undergo a precisely similar change when they are ignited, but the change of these other salts is usually attended by charring.

742. Evolution of GO_2 by Oxidation.—When an oxalate, either in the solid state or in strong solution, is heated with MnO_2 (Note) and H_2SO_4 , it evolves GO_2 :—

 $Am_2C_2O_4 + MnO_2 + 3H_2SO_4 = 2CO_2 + MnSO_4 + 2AmHSO_4 + 2H_2O_5$

Note.—If the MnO_2 , which is to be used in the test, is not known to be free from carbonate, it is best to add a little H_2SO_4 to the MnO_2 in a test-tube and then heat the liquid. If CO_2 is found in the tube (474, 1, 3, 4) the heating should be continued, and the CO_2 should be removed occasionally by sucking fresh air into the test-tube through a glass tube which is pushed down near to the surface of the liquid.

As soon as no CO₂ can be detected in the tube after it has been heated for a short time, the liquid being still distinctly acid, the oxalate is added and the mixture is again heated. CO₂ will be rapidly given off from the oxalate, and may be detected by the ordinary methods.

- 743. KMnO₄-solution, acidified with dilute H₂SO₄, is decolorised in the cold by a soluble oxalate. (Difference from other organic acids.)
- 744. $AgNO_3$ gives a white precipitate of $Ag_2C_2O_4$, which does not darken in colour when the liquid is heated.
- Succinate, $(C_4H_4O_4)''$.—Use Ammonium succinate, $Am_2C_4H_4O_4$. Setemate and benzoate closely resemble one another in their analytical reactions; the differences are therefore important.
- 745. Succinic acid, C₂H₄(COOH)₂, crystallises in monoclinic prisms, which melt at 184° C.: it is readily soluble in water, in alcohol and in ether; but it is insoluble in chloroform. (Difference from benzoic acid.)
- 746. Ignition.—When succinic acid is heated in an ignition-tube it emits fumes of succinic anhydride, which are very irritating to the throat and cause coughing when they are inhaled.

The anhydride condenses as a white sublimate on the cold upper part of the tube.

- 747. KHSO₄, when it is heated with a solid succinate in an ignition-tube, yields a sublimate of succinic acid in oily drops, which crystallise on cooling.
- 748. BaCl₂ solution, when it is added together with AmOH and much alcohol, precipitates white BaC₄H₄O₄. (Difference from benzoate.)

A similar difference is shown with CaCl₂

749. FeCl₃ gives a *reddish-brown* precipitate of basic ferric succinate, which is soluble in mineral acid, but is insoluble in HĀ.

- MALATE, (C₄H₄O₅)".—Use solution of Malic acid, H₂C₄H₄O₅, which has been carefully neutralised by NaOH.
- 750. Malic acid, CH₂:CHOH(COOH)₂, is a colourless, crystalline, deliquescent substance, having a pleasant sour taste: it is readily soluble in water and in alcohol, but not in ether.

When the acid is heated in an ignition-tube it melts at 100° C., and at 180° C. it yields at first malic and fumaric acids; and when it is heated to higher temperatures it chars and evolves CO and CO₂. The crystals have a density of 1.56.

- 751. Strong H₂SO₄, when it is heated with a malate, evolves CO and CO₂, and the liquid gradually darkens.
- 752. CaCl₂: the addition of CaCl₂, followed by boiling or by the addition of alcohol, produces a white precipitate in a neutral solution. This precipitate is soluble in AmOH. (Difference from citrate.)
- 753. $\mathbf{Pb\bar{A}}_2$ gives a white precipitate, which fuses and then dissolves when it is heated with sufficient water. If this precipitate is transferred to a piece of porcelain and heated, it swells up into a light puffy mass.
- 754. AgNO₃ gives a white precipitate, which becomes grey when it is heated.
- /Tartrate, $(C_4H_4O_6)^{\prime\prime}$ or (\bar{T}) .—For a neutral tartrate, use KNa \bar{T} ; for an acid tartrate, $H_2\bar{T}$ or NaH \bar{T} .
 - 755. Tartaric acid, $(CHOH)_2(COOH)_2$, forms colourless rhombic crystals, which are not altered by exposure to the air. It melts at 170° C., has a pleasantly sour taste, and is soluble in water and in alcohol, but not in ether. The acid is detected by being heated either alone or with strong H_2SO_4 (756, 757, 764); and its solution, if neutralized, yields the reactions in paragraphs 758–763.

The normal salts of the alkali-metals are soluble in water, but the acid salts of K and Am are almost insoluble.

- 756. Ignition.—When a solid tartrate is heated strongly in a tube, it chars and gives off a smell of burnt sugar, leaving a residue of the carbonate or the oxide of the metal mixed with carbon.
- 757. Strong $\mathbf{H}_2\mathbf{SO}_4$, when it is heated with a solid tartrate, causes rapid blackening and evolution of a mixture of CO, CO₂ and SO₂.

Accordingly if the mixed gases are passed successively through K₂Cr₂O₇-solution and through lime-water, SO₂ and CO₂ will be detected (474, 489), and the residual stream of CO gas may be kindled and will burn with the characteristic blue flame of CO.

58. AgNO a produces in the strong solution of a neutral tar-

trate a white curdy precipitate of Ag₂T, which is soluble in AmOH and in HNO₃. Keep this precipitate.

759. Production of a Silver mirror.—Allow some of the precipitate from the preceding reaction to settle in a test-tube, and decant as much of the clear liquid as possible.

Then pour very dilute AmOH drop by drop into the tube, until the precipitate, after it has been shaken up with the liquid, is nearly but not quite dissolved.

Now place the test-tube in some cold water in a beaker, and heat the water to about 60° . A brilliant mirror-like film of Ag will be deposited upon the interior of the tube.

Note.—If the original solution of the tartrate is dilute, the mirror is most easily obtained as follows. Very dilute AmOH is added to solution of $AgNO_3$ until the precipitate which forms at first is nearly redissolved; this reagent is added to the neutral tartrate solution, and when the mixture is warmed, the mirror will be formed.

If the liquid is quickly boiled, instead of being gradually heated short of boiling, the Ag will be deposited as a dull black precipitate, which does not adhere to the glass and is not characteristic.

760. CaCl₂ precipitates from the solution of a neutral tartrate white or crystalline CaT: this precipitate often separates only after the liquid has been shaken well, and has then been allowed to stand for a time. It is soluble in acids, and its formation is retarded by the presence of AmCl or of an Am-salt.

When this precipitate is freshly formed, it is completely soluble in strong cold KOH-solution which is free from K_2CO_3 . If this alkaline solution of the tartrate is boiled, the CaT is reprecipitated; but it dissolves again, more or less completely, when the liquid is cold.

Decant the clear liquid from the CaT-precipitate, and pour upon the precipitate a little very dilute AmOH. Then drop in a small crystal of AgNO₃ and heat slowly. A silver mirror will form on the part of the glass on which the crystal of AgNO₃ rests.

761. KCl, or better KĀ, if it is added to the solution of an acid tartrate, or to solution of a neutral tartrate which has been acidified with HĀ, gives a white crystalline precipitate of KHT; this precipitate forms slowly unless the liquid is vigorously stirred or shaken.

The precipitate is soluble in caustic alkalis and in mineral acids, but is almost insoluble in alcohol; hence alcohol is often added to hasten the precipitation.

762. FeCl₃ solution, if it is added in small quantity to solution of a tartrate, undergoes no precipitation when the liquid is made strongly alkaline by the addition of KOH or AmOH.

The precipitation of several other metals, such as Al, Mn and Cu, from their solutions by alkaline hydroxides, is similarly prevented by the presence

* * E .

of a tartrate. This is owing to the formation of a double tartrate, which is not decomposed by the alkali.

763. Hydrogen peroxide.—If a few drops of the peroxide are added to a tartrate solution, then some FeSO₄-solution and excess of KOH, a *violet coloration* will be produced. (Difference from citrate.)

NaOCl-solution, instead of H₂O₂, may be added to the tartrate solution which has been previously acidified with HĀ.

764. Resorcinol.—If a minute quantity of a solid tartrate is mixed with twice its volume of resorcinol and about 2 c.c. of strong $\rm H_2SO_4$ and the mixture gently heated, a bright red coloration is produced. If pyrogallol is substituted for the resorcinol a purple coloration is produced. (Distinction from citrate.) Tartar emetic answers to this test, but not to some of those which are given above.

CITRATE, (C₆H₅O₇)'''.—Use Sodium citrate, Na₃C₆H₅O₇.2H₂O.

765. Citric acid, $C_3H_4(OH)(COOH)_3.H_2O$, crystallises in transparent rhombic prisms. It is very soluble in water, less soluble in alcohol, and only slightly soluble in ether. When the acid is heated to 100° C. it loses its water of crystallisation and becomes anhydrous: it then fuses at 153° C., forms aconitic acid at 175° C., and finally blackens, giving off irritating fumes.

766. A citrate resembles a tartrate and an oxalate in its reactions; it differs by emitting pungent acid fumes when it is carbonised by heat, and by darkening when it is heated with strong H₂SO₄, but only after the mixture has been heated for some time and inflammable gases have escaped.

Citrate differs from tartrate by giving no precipitate with Kā-solution (761), and by yielding no distinctive colours in the reactions in paragraphs 763, 764.

- 767. CaCl₂, or Lime-water.—Calcium citrate is precipitated by the addition of either a large excess of lime-water, or of a mixture of AmOH, AmCl and CaCl₂, but only after long standing or when the liquid is boiled. This precipitate is insoluble in KOH. (Difference from CaT.)
- 768. AgNO₃ gives a white precipitate, which becomes grey when it is heated. Solution of citrate, if it is gently heated with ammoniacal AgNO₃-solution, causes either very slight precipitation of Ag or none at all (758). (Difference from tartrate.)

3. AROMATIC ACID-RADICLES.

The Aromatic Acid-radicles included here are benzoate, salicylate, gallate, tannate, and hippurate. Meconate and urate are also included in this section.

Benzoate, (C7H5O2)'.—Use Ammonium benzoate, AmC7H5O2.

Benzoic acid and benzoates commonly emit an aromatic smell of gum benzoin, from which they have been usually prepared.

769. Benzoic acid, C_6H_5 .COOH, usually crystallises in lustrous flat plates, which melt at 121° and boil at 249° C. It is sparingly soluble in cold water, more soluble in hot water, and readily soluble in alcohol, in ether and in chloroform.

Owing to its slight solubility in cold water the acid is partially precipitated from its solution in alcohol when water is added, or when strong HCl or any other strong acid is added to an aqueous solution of a benzoate.

Most of the benzoates are soluble in water.

770. Ignition.—When benzoic acid is heated, it fuses and evolves dense white fumes, which cause coughing when they are inhaled.

Benzoates give off benzene when they are heated with sodalime: the benzene is recognised by its smell and by burning with a smoky flame, and by the properties described in paragraphs 814 and 815.

- 771. Mineral acids throw down a white crystalline precipitate of benzoic acid, when they are added to the aqueous solution of a benzoate.
- 772. Ethyl benzoate test.—If one c.c. of strong H₂SO₄ and an equal volume of alcohol are poured upon benzoic acid or a benzoate, and the mixture is warmed, a fragrant smell of ethyl benzoate will be perceived.
- 773. FeCl₃ gives a buff-coloured precipitate of basic ferric benzoate which is soluble in AmOH. If this precipitate is heated with strong HCl, benzoic acid separates in crystalline plates. (Difference from succinate.)
- 774. AgNO₃, white precipitate, AgC₇H₅O₂, soluble in hot water, from which it recrystallises on cooling.
- 775. Reduction.—If a little dilute H₂SO₄ and a few pieces of magnesium wire are added to a solution of a benzoate, a fragrant smell of benzaldehyde, C₆H₅COH, will be perceived.

Salicylate, (C₇H₅O₃)'.—Use Sodium salicylate, NaC₇H₅O₃.

- 776. Salicylic acid, C₆H₄.OH.COOH, forms colourless, needle-shaped crystals. The ortho-acid melts at 1.56·5°, and at a somewhat higher temperature it decomposes into CO₂ and phenol or carbolic acid, C₆H₅.OH. It is sparingly soluble in cold water, but more soluble in hot water; it is readily soluble in alcohol and in ether.
- 777. Ignition with Soda-lime.—The decomposition of salicylic acid by heat is rapid and complete, if the acid has been mixed with excess of soda-lime before it is heated; phenol is produced and condenses as a white crystalline sublimate in the cooler part of the tube (848).
- 778. Dilute mineral acids decompose salicylates and cause the separation of white needle-shaped crystals of salicylic acid.
- 779. FeCl₃.—When FeCl₃ is added to solution of salicated it becomes *violet*: the colour is destroyed by HCl, but not by HĀ. (Difference from phenol.)
- 780. Heated with H₂SO₄ and Methyl alcohol, salicylate gives a fragrant smell of methyl salicylate, or "oil of winter-green."
- 781. H_2SO_4 and Nitrite.—The addition to salicylate of strong H_2SO_4 , which has been mixed with a small quantity of 6 per cent. aqueous solution of KNO_2 , causes a *permanent* red coloration. (Difference from phenol.)
- 782. Bromine-water gives a white precipitate of tribrom-salicylic acid.

GALLATE, (C7H5O5)'.—Use Gallic acid, H.C7H5O5.

- 783. Gallic acid, $C_6H_2(OH)_3$.COOH, forms yellowish acicular crystals containing one molecule of water. It is slightly soluble in cold water, readily soluble in hot water, in alcohol and in glycerol, and sparingly soluble in ether.
- 784. Ignition.—When gallic acid is heated, it gives off water at 120° C., melts at about 210° C., and at 253° CO₂ is evolved and pyrogallol (859) sublimes.
- 785. Strong H₂SO₄, when it is heated with gallic acid or a gallate, gives a green solution which changes colour to purple, and SO₂ is eventually evolved.
- 786. FeCl₃ solution yields a bluish-black precipitate which is soluble in excess to a green solution.

- 787. KCN solution, if it is freshly made, gives a red coloration which rapidly disappears, but the colour reappears when the liquid is shaken up with the air. (Distinction from tannate.)
- . 788. Reducing action: when Fehling-solution (909) is warmed with gallic acid, it is gradually reduced with separation of red cuprous oxide. Silver solutions are also reduced by gallic acid with separation of black metallic silver.

TANNATE.—Use tannic acid, C14H10O9.

- 789. Tannic acid in the pure state is a colourless powder, but the commercial acid usually has a yellowish tint. It has a very astringent taste, and is readily soluble in water and in dilute alcohol, but is almost insoluble in absolute alcohol and in ether. It is precipitated when acid is added to its aqueous solution.
- 790. Ignition.—When tannic acid is heated in a tube it blackens, and gives off H₂O, CO₂ and pyrogallol; the pyrogallol condenses in yellowish drops, which crystallise as they cool.
- 791. Strong H_2SO_4 , when it is heated with a tannate, gives a brown coloration, and SO_2 is eventually evolved. (Difference from gallate.)
- 792. FeCl₃ gives a bluish-black precipitate, which is the colouring substance of one of the common inks.
- 793. PbN_2O_6 : white precipitate of lead tannate. (Distinction from gallate and pyrogallol.)
 - 794. KCN solution gives a reddish-brown coloration.
- 795. NaOH dissolves tannic acid, and the solution turns brown in the air by undergoing oxidation.
- 796. Gelatine is precipitated from its solution by tannic acid. (Difference from gallic acid.)

HIPPURATE, $(C_9H_8NO_3)'$.—Dissolve Hippuric acid, $H.C_9H_8NO_3$, in very dilute AmOH-solution, and neutralise when necessary.

^{797.} Hippuric acid, C_6H_5 .CO.NH.CH₂.COOH, occurs in colourless prismatic crystals: it has a bitter taste and a strongly acid reaction: it is only slightly soluble in cold water, but is freely soluble in hot water and in alcohol. The acid melts at 189° C.

^{798.} Ignition with Soda-lime.—When a solid hippurate is heated with soda-lime, it evolves NH_3 and benzene, C_6H_6 (814).

- 799. HCl produces a white crystalline precipitate of hippuric acid in a neutral solution of a hippurate.
- 800. FeCl₃ causes a flesh-coloured precipitate: this is decomposed by HCl with separation of crystalline needles of hippuric acid.
- Meconate, $(C_7H_2O_7)''$.—Use solution of Meconic acid, $H_2C_7H_2O_7$.3 H_2O , neutralising it with dilute AmOH.
- **801.** Meconic acid, $C_5HO_2.OH(COOH)_2$, is present as morphine meconate in opium and in opium extract. It forms colourless crystalline scales, which are slightly soluble in cold water, and readily soluble in hot water and in alcohol.
- 802. Ignition.—When meconic acid is heated in a tube to 100° C. it loses its water of crystallisation, and at 120° C. it forms CO_2 and comenic acid; at a higher temperature it chars and emits the odour of caramel.
- 803. FeCl₃.—Meconic acid may be detected by the production of a purple-brown coloration when it is mixed with FeCl₃-solution. When this liquid is boiled, a precipitate of basic ferric meconate is precipitated, as is the case with ferric acetate or formate.
- 804. PbA_2 solution gives in solution of a meconate a pale-yellow precipitate of lead meconate, which is insoluble in HĀ but soluble in dilute HNO_3 .
- 805. CaCl $_2$ solution gives in neutral solution of a meconate a white precipitate of $CaC_rH_2O_7$.
- 806. AgNO₃ gives in a strong solution of meconate a white precipitate of Ag₂C₇H₂O₂, which changes to bright yellow when a drop of AmOH is added.
 - URATE, (C5H3N4O3)'.—Use Potassium urate, KC5H3N4O3.
- 807. Uric acid, C₅H₄N₄O₃, is a white crystalline powder, which is almost insoluble in cold and hot water, and is insoluble in alcohol and in ether.
- 808. Ignition.—When a urate is heated in an ignition-tube it chars and evolves NH₃, HCNO and HCN. If it has been mixed with NaOH or with soda-lime before being heated, NH₃ is evolved and NaCN is left in the solid residue.
- **809.** $\mathbf{HNO_3}$ dissolves the acid with effervescence: if this solution is evaporated to dryness in a porcelain dish, and the residue is then moistened with AmOH, unless AmOH was originally present, bright red *murexide* is formed; this becomes purple when KOH is added.
 - 810. HCl precipitates white uric acid from solution of a urate.

GROUP II.—HYDROCARBONS AND THEIR DERIVATIVES.

• The substances included under this heading are turpentine, benzene, naphthalene, anthracene, chloroform, carbon tetrachloride, iodoform and carbon disulphide.

TURPENTINE, C10H16.

811. Turpentine is a mixture of a number of isomeric liquids which have the formula $C_{10}H_{16}$, pinene being the constituent which is present in largest proportion. Turpentine possesses a characteristic smell; it boils at $160\,^{\circ}$ C., and its density is 0.876. The warm liquid is easily kindled, and burns with a very smoky flame. It is not miscible with water, but readily dissolves in strong alcohol and in ether.

812. Strong H₂SO₄, when it is mixed in small quantity with turpentine, causes a peculiar and very characteristic smell to be evolved when the mixture is warmed.

813. Br-water, when it is shaken up with turpentine, is at once decolorised.

BENZENE, C.H.

814. Benzene is a very mobile, volatile, strongly smelling liquid which burns readily with a smoky flame. It boils at 80 ° C., and its specific gravity is 0.8839. It is miscible with alcohol and with ether, but is insoluble in water.

815. HNO_3 .—If strong fuming nitric acid, or a mixture of strong H_2SO_4 and strong HNO_3 , is warmed with benzene, nitrobenzene, $C_6H_5NO_2$, is produced: and this substance separates as a reddish-yellow oil, smelling of bitter almonds, when the acid liquid is largely diluted with water.

If this nitro-benzene is poured into dilute HCl and pieces of zinc are introduced, aniline is formed and may be detected by the tests in paragraphs 980-982.

NAPHTHALENE, C₁₀H₈.

816. Naphthalene crystallises in shining plates, which melt at 80° C., and when further heated boil at 218° C. Naphthalene is insoluble in water, but is readily soluble in hot alcohol and in ether. It possesses a characteristic smell and is very volatile, the vapour

burning with a bright smoky flame. Naphthalene vapour is always present in coal-gas, and contributes largely to the illuminating power of the gas.

ANTHRACENE, C14H10.

817. Anthracene crystallises in colourless glistening leaflets, which show a fine blue fluorescence. It melts at 216.6°, and boils at 351° C. It dissolves readily in hot benzene, but sparingly in alcohol and in ether.

CHLOROFORM, CHCl3.

- 818. Chloroform is a colourless, mobile liquid, with an ethereal smell and sweetish taste, which boils at 61.2°, and has a density of 1.5. It is only slightly soluble in water, but is miscible with alcohol and with ether. Its vapour, when inhaled, acts as a powerful anæsthetic.
- 819. Formation of Phenyl isocyanide.—When chloroform is mixed with a drop of aniline, and with some alcoholic solution of KOH, and the mixture is heated, an *intensely disagreeable* smell of phenyl isocyanide is given off.
- 820. Fehling's-solution (909), when it is warmed with chloroform, deposits red cuprous oxide after some time.
- 821. Phenol and KOH.—If a mixture of alcoholic solution of phenol with KOH is evaporated to dryness on the water-bath, the residue assumes a beautiful purple colour when it is moistened with chloroform. (Difference from chloral.)

CARBON TETRACHLORIDE, CCl₄.

- 822. Carbon tetrachloride is a mobile liquid, with a boiling-point of 76.7° and a density of 1.6. It is insoluble in water but miscible with alcohol and with ether.
- 823. Phenyl isocyanide test.—When carbon tetrachloride is heated with a drop of aniline and with alcoholic potash, the nauseous smell of phenyl isocyanide is produced, but more slowly than with chloroform (819).

IODOFORM, CHI₃.

824. Iodoform is a bright yellow substance, which emits a characteristic smell, and crystallises in hexagonal plates or stars. It

has a density of 4-01, melts at 119° C., and when further heated it vaporises and evolves iodine and HI, leaving a residue of carbon. It is insoluble in water and in acids, but dissolves in alcohol and in ether.

When it is boiled with KOH-solution, iodoform breaks up into a mixture of potassium formate and potassium iodide, which may be separately identified (728-730 and 552-556).

When iodoform is warmed with solution of phenol in alcoholic NaOH, a red coloration is produced.

CARBON DISULPHIDE, CS2.

- 825. Carbon disulphide is a mobile, highly refractive liquid. When it is pure it possesses a pleasant ethereal smell; but it usually emits an extremely repulsive odour, which is due to the presence of impurity. It boils at 46° C., and has a density of 1·292. It is not miscible with water, but dissolves in alcohol and in ether; it readily dissolves fatty bodies and oils. CS_2 is volatile, and the vapour is readily kindled, burning with a blue flame and producing SO_2 and CO_2 .
- 826. Formation of Potassium xanthate.—If CS₂ is heated with alcoholic solution of KOH, potassium xanthate, CS.OC₂H₅.SK, is formed: the addition of CuSO₄-solution to this liquid produces a yellow precipitate of copper xanthate.
 - 827. **KOH**, when it is boiled with CS_2 , forms a brown solution containing K_2CO_3 and K_2CS_3 : on the addition of one drop of $Pb\bar{A}_2$ -solution, black PbS is precipitated.

NITROBENZENE, C₆H₅NO₂.

- 828. Nitrobenzene is a pale-yellow liquid, smelling of bitter almonds: its density is 1.187, and it boils at 210.9° : it is insoluble in water, but is miscible with most organic liquids.
- 829. Reduction to aniline: if a lit! e nitrobenzene is mixed with dilute HCl and a few small pieces of zinc are introduced, aniline is gradually formed, and may be detected as is described in paragraphs 980-982.

GROUP III.—ALCOHOLS AND ETHERS.

In this Group there are included methyl alcohol, ethyl alcohol, amyl alcohol, glycerol, ethyl ether and ethyl acetate.

METHYL ALCOHOL, CH₃.OH.

- 830. Methyl alcohol resembles ethyl alcohol (834) in its general properties. It boils at 64.7° C., and has a density of 0.81. It is miscible with water and with ether in all proportions. Its vapour, when it is kindled, burns with a pale blue flame.
- 831. Oxidation to Formic acid.—If a mixture of methyl alcohol with dilute H₂SO₄ and K₂Cr₂O₇ is distilled (32, 672), the distillate will contain formic acid; and when this acid distillate is warmed with AgNO₃, it will cause a brilliant silver mirror to be deposited.

When ethyl alcohol is oxidised by similar treatment, it yields aldehyde, and this only deposits a silver mirror when it is warmed with AgNO₃ in an alkaline solution (879).

- 832. Formation of Methyl salicylate.—When methyl alcohol is heated with strong $\rm H_2SO_4$ and salicylic acid, a fragrant smell of methyl salicylate, or "oil of winter-green," is evolved.
- 833. Iodoform is not produced when pure methyl alcohol is warmed with excess of iodine in the presence of $\rm Na_2CO_3$ (836). (Difference from ethyl alcohol.)

ETHYL ALCOHOL, C₂H₅.OH.—Use rectified spirit diluted with four times its volume of water.

834. Ethyl alcohol is a mobile fragrant liquid: it boils at 78°, has a density of 0.7937 at 15° C., and its vapour burns with a pale blue smokeless flame. It is miscible in all proportions with water and with ether.

If much water is prese, with the alcohol, it may be necessary to add dry K_2CO_3 in excess to the liquid, and then to distil over about one-third of it (32,672). The alcohol is thus concentrated, and is much more readily detected by the following tests in the distillate than in the original dilute liquid.

- 835. Strong H₂SO₄, when it is mixed with alcohol and heated, gives off ether and ethylene, which burn with a *luminous* flame.
- 836. Formation of Iodoform.—If Na₂CO₃-solution is added to water which contains a little alcohol, and the mixture is gently

heated for some time, small pieces of iodine being occasionally added until the solution retains a brown colour, golden yellow crystals of iodoform will separate. If very little alcohol was present, the liquid may require to stand for ten or twelve hours before the crystals make their appearance.

When these crystals are examined under the microscope, they are seen to be hexagonal tablets or six-pointed stars.

This reaction is yielded by other substances, such as aldehyde, acetone and ethyl acetate, and is therefore not altogether characteristic of ethyl alcohol.

837. Formation of Aldehyde and Acetic acid.—Alcohol may be converted by oxidation into aldehyde and acetic acid, by heating it for some time with K₂Cr₂O₇ and H₂SO₄. During the process of oxidation, aldehyde will be first smelt, and then acetic acid.

If the smell is not distinctly recognised, the liquid may be distilled and the first portion of the distillate divided into two portions: one portion may then be tested for aldehyde by paragraph 880, and the other may be neutralised with Na₂CO₃, evaporated to dryness, and the solid residue tested for acetate (733-737).

It must be remembered that acetic acid may result from the oxidation of other substances besides alcohol.

838. Benzoic- and Acetic-ether.—When ethyl alcohol is heated with strong H₂SO₄ and a benzoate or an acetate, the characteristic smell of ethyl benzoate or of ethyl acetate will be recognised.

AMYL ALCOHOL, C₅H₁₁.OH.

- 839. The commercial alcohol is isoamyl alcohol; it is a colourless oily liquid, and its vapour has a strong smell and causes coughing when it is inhaled. The alcohol is only slightly soluble in water; it boils at 131.6° C., and has a density of 0.82.
- 840. Strong H₂SO₄, when it is warmed with amyl alcohol, gives a reddish-brown coloration: if a few drops of glacial acetic acid and one c.c. of FeCl₃-solution are then added, a purple colour is produced in the warm liquid.
- 841. Sodium acetate, if it is added together with a little dilute H₂SO₄ and warmed, gives a pear-like odour, which is due to the formation of amyl acetate.

GLYCEROL, OR GLYCERINE, C3H5(OH)3.

842. Glycerol is a colourless, hygroscopic, viscous liquid, with an intensely sweet taste. It has a density of 1.27, a melting point

of 20° C., and a boiling-point of 290°. It burns with a somewhat luminous flame, and is miscible with water and with alcohol, but not with ether.

- 843. NaHSO₄.—When glycerine is heated with NaHSO₄, ît evolves intensely pungent acrolein vapour. If this vapour is passed into solution of rosaniline, which has been previously decolorised by SO₂, it produces a red coloration. Other aldehydes produce a similar coloration.
- 844. Phenol and H₂SO₄.—If phenol, strong sulphuric acid and glycerine are mixed together in equal volumes, and the mixture is heated to 120° C., and is then diluted and mixed with excess of AmOH, it becomes *crimson* in colour.

ETHYL ETHER, (C₂H₅)₂O.

845. Ether is a very mobile, volatile, fragrant liquid, which is easily kindled and burns with a bright flame. It has a boiling-point of 34.6° C., and a density of 0.7201. Ether dissolves in about ten times its volume of water, and is miscible with alcohol and with other organic liquids.

ETHYL ACETATE, OR ACETIC ETHER, C2H3C2H3C2.

- 846. Ethyl acetate, CH₃.COOC₂H₅, is a very mobile, fragrant liquid. It has a boiling-point of 77·2° C., and a density of 0·9048: it is somewhat soluble in water, and is readily soluble in alcohol.
- 847. KOH.—Acetic ether is hydrolysed by KOH in a manner which is typical of similar organic bodies:—

 $C_2H_5.C_2H_3O_2 + KOH = C_2H_5.OH + KC_2H_3O_2$

This change may be effected by simply warming the liquid for some time with KOH.

When the resulting liquid is distilled, alcohol may be detected in the distillate (836), and acetate may be found in the distillation flask (733-737).

GROUP IV.—PHENOL AND ITS DERIVATIVES.

PHENOL, OR CARBOLIC ACID, C₆H₅.OH.—Use solution of a small crystal of the substance in hot water.

- 848. Phenol is a colourless crystalline substance; the commercial substance, however, usually has a pinkish tinge. Its smell is very characteristic. It melts at 43° C. and boils at 181° C., and has a density of 1.05. It is slightly soluble in water, but freely soluble in benzene, in alcohol and in ether. It is also soluble in caustic alkali solution, but insoluble in solution of alkaline carbonate. (Distinction from the true acids.)
- 849. FeCl₃ solution gives with phenol solution a violet coloration, which is destroyed by $H\bar{A}$.
- 850. Bromine-water, or Bleaching-powder solution, when it is added to phenol solution, which has been previously mixed with a quarter of its volume of dilute AmOH, produces a blue coloration: this colour changes to red when an acid is added.
- 851. Liebermann's reaction.—If a crystal of KNO_2 is dissolved in strong $\mathrm{H}_2\mathrm{SO}_4$ and the solution is added to phenol, a green coloration is produced, or a brown coloration if sufficient acid is not present. If this solution is then poured into water a red coloration is produced, and this changes back to green when the liquid is made alkaline by adding NaOH.
- **852.** Bromine water, when it is added to phenolesolution, gives a pale yellow precipitate of tribromphenol bromide, $C_6H_2Br_3.Br$.
- 853. Strong HNO $_3$, where it is warmed with a little phenol solution, gives a bright yellow liquid containing picric acid: the colour of this solution is made more intense by adding NaOH. If this solution is boiled with a little glucose its colour changes to deep red, owing to the formation of picraminic acid.

Hydroquinone, $C_6H_4(OH)_2$.

- 854. Hydroquinone is a white crystalline substance which shows a greenish tint. It melts at 170.3°, boils at 285° C., and is soluble in water, in alcohol, and in ether, but is insoluble in benzene.
- 855. FeCl₃, if it is added to a dilute solution of hydroquinone, gives a green coloration which rapidly changes to reddish-brown. If FeCl₃ is added to a strong solution, or to solid hydroquinone, a dark green crystalline precipitate is formed.
- 856. AgNO₃ solution is reduced by hydroquinone solution, and metallic Ag is precipitated.
- 857. KOH solution becomes brown when it is shaken with hydroquinone solution.
- 858. KMnO₄ solution acidified with dilute H₂SO₄, when it is boiled with hydroquinone solution, evolves the irritating odour of quinone.

Q, A,

Pyrogallol, or Pyrogallic Acid, C6H3(OH)3.

- 859. Pyrogallol is a white crystalline substance: it melts at 132° C., boils at 293°, and is readily soluble in water, in alcohol and in ether, but is less soluble in chloroform and in benzene.
- 860. KOH and other caustic alkalis readily dissolve this substance, and the solutions rapidly become brown by absorption of oxygen from the air.
 - 261. FeSO4 colours pyrogallol solution dark blue.
 - 862. FeCl₃ gives a green coloration to pyrogallol solution.
- 863. AgNO₃ is immediately reduced by pyrogallol solution, and metallic silver is precipitated.
 - 864. Ca(OH), gives a purple coloration which changes to brown.
- 865. Fehling's solution (909) is reduced when it is heated with pyrogallol solution, and red Cu₂O is precipitated.

TRINITROPHENOL, OR PICRIC ACID, C6H2(NO3)3.OH.

- 866. Pieric acid is a yellow crystalline substance, which melts at 122.5°C. Its taste is intensely bitter and it stains the skin yellow. It is very slightly soluble in cold water, but is readily soluble in hot water, in alcohol, and in ether. The metallic pierates undergo violent explosive decomposition when they are placed under suitable conditions.
- 867. Ignition: picric acid fuses when it is heated in an ignition-tube, and then explodes slightly, giving off dense black furnes.
- 868. Bleaching-powder solution, when it is heated with picric acid, gives off the pungent odour of chlorpicrin, which resembles that of mustard-oil.
- 869. KOH and Glucose, when they are heated with pieric acid, yield an intense red colour, which is due to the formation of picraminic acid.
- 870. Ammonia-copper sulphate solution, made alkaline with AmOH, gives a bright green precipitate.
- 871. NaOH dissolved in alcohol gives a deep yellow precipitate of sodium picrate, $C_6H_2(NO_3)_3ONa$.

GROUP V.—ALDEHYDES AND KETONES.

FORMALDEHYDE, H.COH.—Use, "formalin" liquid.

872. Formaldehyde is a gas with a peculiar pungent smell.

in water. When the aqueous solution of formaldehyde is evaporated on the water-bath, white solid paraformaldehyde is produced.

- 873. Ammoniacal silver nitrate (879), when it is mixed in a test-tube with a little formaldehyde, and the mixture is warmed by placing the tube in a beaker of hot water, yields a brilliant silver coating on the inside of the test-tube :—H.COH + $Ag_2O = H.COOH + Ag_2$.
- 874. Fehling's solution (909) is reduced when it is warmed with formaldehyde, and red cuprous oxide is precipitated. (Aromatic aldehydes do not reduce Fehling's solution.)
- 875. Schiff's reaction.—If formaldehyde is added to a dilute solution of magenta (fuchsine), which has been previously decolorised by means of sulphurous acid, the solution will gradually become pink and ultimately magenta in colour.
- 876. Pyrogallol in freshly prepared solution, if it is added to formaldehyde and the mixture is then acidified with *strong* HCl, yields gradually a white precipitate which eventually changes in colour to deep red.
- 877. Phenylhydrazine hydrochloride solution, if it is mixed with an equal volume of dilute sodium nitroprusside solution, and is then added to a little dilute formaldehyde solution and rendered alkaline with NaOH, gives an intense blue coloration.

ACETALDEHYDE, CH₃.COH.

- 878. Acetaldehyde is a very mobile liquid, possessing a most characteristic smell: it boils at 20.8° C., and has a density of 0.778, unless it is partially polymerised. It is miscible with water, with alcohol and with ether in all proportions.
- 879. AgNO₃ solution, if it is mixed with very dilute AmOH until the precipitate which forms at first just disappears, and is then heated with acetaldehyde, produces a silver mirror on the glass. The test is made as is directed in paragraph 759.
- 880. Schiff's reaction.—Magenta (fuchsine) solution, which has been decolorised by SO₂, produces a violet-red colour when it is mixed with acetaldehyde.
- 881. KOH solution, when it is heated with acetaldehyde, produces aldehyde-resin, a yellow substance with a peculiar smell.
 - 882. H2S, when it is passed into the aqueous solution of alde-

hyde, produces an oily liquid, which is changed by acids into a solid polymer.

- 883. NaHSO₃, in saturated solution, gives a crystalline precipitate (C₂H₄O.NaHSO₃), which is decomposed by acids and by alkalis with liberation of aldehyde. This reaction is common to many aldehydes.
- 884. Pyrogallol and strong HCl (876) give with acetaldehyde a white precipitate, which gradually turns yellow. With a dilute solution the precipitate only appears after a time.

Chloral, CCl₃.COH.—Use Chloral hydrate, CCl₃.CH(OH)₂, dissolved in water.

- 885. Chloral is a colourless liquid, with a pungent tear-exciting smell: it boils at 97.7°, and has a density of 1.512. It unites with water, forming crystalline chloral hydrate, which melts at 47.4°, boils at 97.5°, and has a density of 1.58. Choral readily dissolves in water, in alcohol and in ether.
- 886. Phenyl isocyanide test.—When chloral hydrate is heated with alcoholic solution of KOH and aniline, it yields phenyl isocyanide, which is recognised by its extremely offensive smell (819).
- 887. Silver mirror.—When chloral hydrate is warmed in a test-tube with ammoniacal AgNO₃-solution (759) and a drop of KOH solution, a silver mirror is readily deposited on the inside of the tube. (Difference from chloroform.)
- 888. Fehling's solution (909) is reduced when it is warmed with chloral hydrate, and red Cu₂O is formed.
- 889. Schiff's reaction.—Chloral gives the magenta reaction (875), whereas chloral hydrate does not.
- 890. KOH.—Chloral hydrate is decomposed by aqueous KOH solution: chloroform separates in minute drops, and potassium formate remains in solution.

Benzaldehyde, C₆H₅,COH.

- 891. Benzaldehyde is a colourless and highly refractive liquid with a fragrant smell: it has a density of 1.05, and boils at 179° C.: it is sparingly soluble in water, but readily soluble in alcohol and in ether. On exposure to the air it is slowly converted into benzoic acid (769 et seq.).
- 892. KOH: when benzaldehyde is heated with solid KOH, benzyl alcohol and potassium benzoate are formed; and if the resulting solution is acidified with HOL, benzoic acid is procysitated (771).

893. AgNO $_3$ in ammoniacal solution is reduced by this aldehyde to metallic silver.

894. Na₂SO₃ solution dissolves benzaldehyde: the careful addition of dilute H₂SO₄ to this solution precipitates crystalline flocks of C₇H₆O.NaHSO₃. Schiff's reaction (875) is given by benzaldehyde.

ACETONE, OR DIMETHYL-KETONE, CO(CH₃)₂.

- **895.** Acetone is a mobile liquid, with a peculiar and characteristic smell: it boils at $56\cdot1^{\circ}$ C., and has a density of $0\cdot7965$: it is miscible with water, alcohol and ether in all proportions.
- 896. Iodoform reaction: if a few drops of solution of iodine in KI-solution are mixed with solution of acetone, and Na₂CO₃-solution is then added until the brown colour of the mixture disappears, and the liquid is warmed, iodoform will separate in golden yellow plates as the liquid cools.

This reaction is also given by ethyl alcohol.

- If, however, a solution of acetone is made alkaline with AmOH and a solution of I in KI-solution is added drop by drop, a black precipitate of nitrogen iodide will form: this black precipitate will disappear when the liquid is gently warmed, and iodoform will then crystallise out when the liquid is cooled. Ethyl alcohol does not yield iodoform under these conditions.
- 897. HgCl₂ solution, if it is made strongly alkaline by alcoholic solution of KOH, yields a precipitate of HgO: when the liquid containing the precipitate is shaken with acetone, part of the HgO is dissolved; if the liquid is then filtered, Hg may be shown to be present in the filtrate either by adding Am₂S (134), or by acidifying the liquid with HCl and adding SnCl₂ (135).
- 898. Sodium nitroprusside, if it is added to acetone solution which has been previously mixed with twice its volume of strong NaOH-solution, produces a *bright red* coloration: this coloration becomes bluish when HA is added.
- 899. NaHSO₃, in saturated solution, gives a crystalline precipitate (C₃H₆O.NaHSO₃), which is decomposed by acid or alkali with liberation of acetone.
- 900. Parabromphenylhydrazine, if it is dissolved in glacial acetic acid and is then diluted and mixed with acetone, gradually yields yellowish needles of bromphenyl hydrazone.

GROUP VI.—CARBOHYDRATES.

gor. The general name of carbohydrate has been given to those compounds of carbon, hydrogen and oxygen which contain hydrogen and oxygen in the atomic relation of $H_{2n}O_n$, that is in the proportion in which these elements are present in water.

This Section gives the reactions of the sugars; and includes the glucoses, dextrose and levulose ($C_6H_{12}O_6$); and the saccharoses, sucrose, maltose and lactose ($C_{12}H_{22}O_{11}$). The reactions of carbohydrates of the general formula ($C_6H_{10}O_5$)_n, including cellulose, starch and dextrin, are also given.

GENERAL REACTION FOR THE SUGARS.

902. When a sugar, which contains an aldehydic group (COH) or a ketonic group (CO), is heated with one molecule of phenyl hydrazine, it is converted into a hydrazone: thus glucose forms glucose phenyl hydrazone:—

Glucose. Phenyl hydrazine. $CH_2(OH)(CH.OH)_4COH + C_6H_5NH.NH_2 = Glucose$ phenyl hydrazone. $CH_2(OH)(CH.OH)_4CH : N.NHC_6H_5 + H_2O$

If an excess of phenyl hydrazine is added, the final CHOH group is

oxidised to CO, and part of the phenyl hydrazine is reduced to aniline and NH₃: $-C_6H_5NH.NH_2 + H_2 = C_6H_5NH_2 + NH_3$.

The oxidation product of the phenyl hydrazine then combines with a second molecule of phenyl hydrazine and yields an osazone:—

 $CH_2(OH)(CH.OH)_3CO.CH : N.NH.C_6H_5 + C_6H_5.NH.NH_2 =$

Osazone. $CH_2(OH)(CHOH)_3: C(N.NHC_0H_5).CH: N.NH.C_0H_5 + H_2O$

The melting-point of the osazone is characteristic of the sugar from which it has been derived. Hence the preparation of an osazone (903) is often carried out in order to identify a sugar.

903. Preparation of Osazones.—Dissolve half a gramme of the sugar in 5 c.c. of water in a test-tube, add 3 c.c. of phenyl hydrazine and then 3 c.c. of acetic acid, and place the test-tube in boiling water for about ten minutes. The osazone will separate in yellow crystals, which are filtered off, washed with water, and dried. Their melting-point is then determined (669, 670).

DEXTROSE, DEXTRO-GLUCOSE, OR GRAPE SUGAR, C. H12O.

^{. 904.} Dextrose is a colourless soluble solid, with a sweet taste. It crystallises from its solution in alcohol in the anhydrous state,

and these crystals fuse at 146°C .: from its solution in water it crystallises in a hydrated form, $C_6H_{12}O_6.H_2O$, which melts at 86°C . Dextrose is readily soluble in water and in dilute alcohol, but is insoluble in strong alcohol and in ether. Its density is 1.55, and its specific rotatory power is $[a]_D + 52.8^{\circ}\text{C}$. Dextrose can be fermented by yeast. Its osazone melts at 204° .

- 905. Ignition.—When glucose is heated in an ignition-tube, it melts at about 86°C., becomes anhydrous at 146°C., and turns brown at higher temperatures giving off the characteristic smell of burnt sugar.
- 906. Strong H₂SO₄.—Dextrose is distinguished from other sugars, by yielding only a *yellow liquid* when strong H₂SO₄ is added to its strong solution, and the mixture is kept cool. (Distinction from cane-sugar.)

When this yellow liquid is heated, it blackens.

- 907. KOH.—When dextrose is warmed with strong KOH-solution, the liquid becomes brown. (Distinction from cane-sugar.)
- 908. Reduction of CuO.—If a few drops of CuSO₄-solution are mixed with dextrose-solution, and excess of KOH-solution is then added, no precipitate is formed; but when this mixture is heated, yellow cuprous hydroxide is formed, and this is converted into red Cu₂O when the liquid is more strongly heated.

The dextrose solution may be advantageously mixed with Fehling-solution (909) in this test, instead of being mixed with CuSO₄ and KOH.

- 909. Fehling solution is prepared as follows: 35 grammes of crystallised ${\rm CuSO_4.5H_2O}$ are dissolved in one litre of water, and the clear solution is kept in a stoppered bottle. 173 grammes of Rochelle-salt (KNaT), and sixty grammes of NaOH are then dissolved in a litre of water; this solution is kept in a corked bottle. Equal volumes of these two solutions are mixed together in order to prepare the Fehling-solution.
- 910. Silver mirror.—If dextrose is gently heated with ammoniacal AgNO₃-solution (759, *Note*) in a glass vessel, a silver mirror will be produced on the interior of the glass.
- 911. Other Reduction tests.—Dextrose differs from maltose and lactose by forming red Cu₂O when it is boiled with CuĀ₂-solution containing HĀ. When dextrose is mixed with solutions of KOH and of basic bismuth nitrate and heated, it yields a black precipitate.
- 912. Picric acid.—If dextrose is boiled with KOH-solution and a little picric acid a yellow colour is produced, which

changes to deep red: the colour is due to the formation of picraminic acid:

913. Fermentation.—If clean freshly-washed yeast is added to dextrose-solution in the flask shown in Fig. 56, page 135, and the temperature of the liquid is maintained at about 25°C for an hour or more, fermentation will take place with the production of ethyl alcohol and carbon dioxide gas.

The escaping gas may be shown to be CO₂ by letting it bubble through lime-water (474, 4), and ethyl alcohol may be found in the liquid which remains in the flask by the reactions in paragraphs 836, 837.

This reaction by fermentation is given by other sugars.

- 914. Osazone.—When dextrose is heated with phenyl hydrazine and acetic acid (903) yellow crystals of the osazone are formed, which melt at 204°C.
- 915. Lead acetate, if it is added to glucose solution and the liquid is boiled, then mixed with AmOH in quantity just sufficient to produce a white precipitate and further boiled, yields a pink precipitate. (Difference from sucrose and lactose.)

LEVULOSE, OR FRUCTOSE, C₆H₁₂O₆.

916. Levulose is usually a colourless, difficultly crystallisable syrup. It is sweeter than dextrose, and is also more soluble in water and in alcohol. Its specific rotatory power is $[\alpha]_D - 102^\circ - 0.56.t^\circ$, and its melting-point is 95°C. It can be fermented by yeast, but differs from dextrose by forming an insoluble compound with lime. Its osazone (903) melts at 204°C.

Sucrose, or Cane-sugar, C₁₂H₂₂O₁₁.

- 917. Cane-sugar is a colourless crystalline substance, which is easily soluble in water and is intensely sweet in taste. It is only sparingly soluble in alcohol. Cane-sugar fuses at $160\,^{\circ}$ C, and is readily charred by heat. Its density is 1.59, and its specific rotatory power is $[a]_D + 66.5^{\circ}$. It is not readily fermented by yeast, and it does not form an osazone.
- 918. Strong H₂SO₄ produces a deep brown coloration if the mixture is kept cool; but the liquid ultimately becomes black owing to the separation of carbon. (Difference from other sugars.)

 The presence of much water prevents this change from occurring

at ordinary temperatures, but the liquid darkens when it is heated.

- 919. KOH produces no coloration (907), and neither silver solution nor alkaline copper-solution is reduced (910, 912). (Difference from dextrose, maltose and lactose.)
- 920. Conversion into Glucose.—Sucrose solution, when it is heated for a minute with a few drops of strong HCl, yields a mixture of equal quantities of dextrose and levulose which is known as "invert sugar." After this acid sugar-solution has been neutralised by KOH, it reduces Fehling-solution (909).
- 921. Citric acid, in 5 per cent. solution, when it is boiled with sucrose also hydrolyses it and forms glucoses. (Distinction from maltose and lactose.)
- 922. Cane-sugar may be identified in the presence of dextrose maltose and lactose by its inability to reduce Fehling-solution (909).

If a solution which contains these sugars is mixed with Fehling-solution and heated on a water-bath until no further reduction takes place, as is shown by the blue colour of the liquid remaining, any sugar still in solution will be sucrose. The precipitated Cu₂O may be filtered off and the filtrate acidified with dilute H₂SO₄ and boiled for five minutes, then rendered alkaline with NaOH and again heated with Fehling-solution: reductior and separation of Cu₂O will take place if cane-sugar was present since it has been converted into "invert sugar" by being heated with the acid (920).

MALTOSE, OR MALT-SUGAR, C12H22O11.H2O.

- 923. Maltose is a sweet sugar, with a density of 1.54: it is very soluble in water, but is almost insoluble in absolute alcohol. It resembles dextrose in many of its properties, and is readily fermented by yeast. Its specific rotatory power is $[a]_D + 137^\circ$ and its osazone fuses at 206°C .
- 924. Ignition.—Maltose loses water when it is heated, and is ultimately decomposed with charring.
 - 925. Strong H₂SO₄ forms a clear solution with maltose in the cold, but this gradually carbonises and blackens when it is heated
 - 926. NaOH solution, when it is boiled with maltose, become yellow and finally brown.
 - 927. Fehling solution (909) is reduced by maltose.
 - 928. CuA₂-solution, mixed with HĀ, is not reduced by maltos on boiling (Distinction from glucose.)

- 929. Lead acetate, if it is boiled with maltose solution, and AmOH is then added until a precipitate is produced and the solution is again boiled, yields a precipitate which is finally pink in colour.
- 930. Picric acid, mixed with excess of KOH, yields with maltose the same result as it does with glucose (912).
- 931. Dilute Sulphuric acid, when it is boiled with maltose solution, hydrolyses it, yielding two molecules of dextrose.
- 932. Osazone.—When maltose is heated with $C_6H_5NH.NH_2$ and $H\bar{A}$ (903) a yellow crystalline osazone is produced, which has a melting-point of 206°C.

LACTOSE, OR MILK-SUGAR, C12H22O11.H2O.

- 933. Lactose is much less soluble in water, and less sweet in taste, than the sugars which have been already described, and is insoluble in absolute alcohol and in ether. Lactose melts at 203.5° C., has a density of 1.52, and is not fermented by pure yeast: its specific rotatory power is $[a]_D + 52.5^{\circ}$, and its osazone melts at 200° C.
- 934. Ignition.—When lactose is heated, water is given off, and charring gradually takes place.
- 935. Strong H₂SO₄ produces no change until heat is applied, when blackening is gradually produced, but less readily than with cane-sugar.
- 936. NaOH solution produces a yellow solution when the liquid is boiled, and the colour gradually darkens to brown.
- 937. Fehling-solution (909) and ammoniacal AgNO₃ (759, *Note*) are each of them reduced by lactose.
- 938. CuÃ₂ and HĀ: no reduction is produced by lactose on boiling, unless the solution is very strong.
- 939. Lead acetate, when it is boiled with lactose solution, and sufficient AmOH is then added to produce a white precipitate, and the liquid is again boiled, produces a yellow precipitate.
- 940. Picric acid, mixed with excess of KOH, behaves with lactose as it does with glucose (912).
- 941. Osazone.—When lactose is heated with C₆H₅.NH.NH₂ and HĀ, a yellow crystalline osazone is produced, with a meltingpoint of 200°.

Test.	Glucose.	Sucrose.	Lactose.	Maltose.
Heat in ignition- tube.	Melts, and chars when it is fur- ther heated.	Melts, and chars readily.	Chars without previously melting.	Melts, and chars when it is fur- ther heated.
Add strong H ₂ SO ₄ and heat.	Chars gradually when it is heated.	Chars in the cold.		Chars slowly
Heat with NaOH- solution.	Becomes brown.	Does not change.	Becomes brown.	Becomes brown.
Add Pb \$\overline{\Lambda}_2\$-solution and AmOH, and heat gently.	Pink precipitate.	White precipi-	Yellow precipitate.	Pink precipitate.
Add CuÃ2-solu- tion and HÃ and heat.	Cu ₂ O separates on boiling.	No change.	No change.	No change.
Add Fehling-solu- tion and heat.	Cu ₂ O separates on boiling.	No change.	Cu_2O separates on boiling.	Cu_2O separates on boiling.
Add Phenyl hydrazine and HA (903) and heat.	Yellow crystals of glucosazone, M.P. 206°.	No osazone.	Yellow crystals of lactosazone, M.P. 200°.	Yellow crystals of maltosazone, M.P. 206°,
Specific rotatory power.	$[a]_{\mathbf{D}} = +52.8^{\circ}.$	$[a]_D = +66.5^{\circ}.$	$[a]_{D} = +52.5^{\circ}.$	$[a]_{D} = +137^{\circ}.$

942. SUMMARY OF REACTIONS OF THE SUGARS.

CELLULOSE, (C₆H₁₀O₅)_n.—Use Cotton-wool.

- 943. Cellulose is a colourless, insoluble, tasteless substance: it is unaffected by water and by most other solvents, even when it is boiled with them.
- 944. Strong H₂SO₄ dissolves cellulose, and when the solution is diluted and boiled, the cellulose is hydrolysed and changed completely into dextrose; after this solution has been neutralised, the dextrose may be detected by means of Fehling-solution (908).
- 945. Iodine reaction.—Cellulose itself gives no reaction with iodine; but after it has been treated with one of the following mixtures (a, b) iodine produces a blue colour with the product, which is known as "amyloid."
- (a) A mixture of 90 parts of $ZnCl_2$ -solution of $2\cdot 0$ specific gravity, with 6 parts of KI and 10 parts of water.
- (b) A mixture of 3 parts of strong H₂SO₄ with 2 parts of glycerol and 1 part of water. This mixture is rubbed with the cellulose in order to effect the necessary change.
- 946. Cuprammonium hydroxide, made by dissolving freshly-precipitated Cu(OH)₂ in *strong* ammonia-solution, dissolves cellulose: the cellulose is reprecipitated when an acid is added in excess to the solution.

STARCH.—Use Potato- or Rice-starch in powder, (C₆H₁₀O₅)_n.

- 947. Starch is insoluble in cold water, but it yields a transparent liquid when it is *boiled* with water; this liquid becomes gelatinous as it cools, if much starch is present.
- 948. Starch granules may be identified by their appearance, when they are examined by a quarter-inch object-glass under the microscope. If they are viewed as transparent objects they usually appear round or oval, but they are occasionally irregular in shape, and show dark rings which are more or less concentric. Many kinds of starch-granules show a cross when they are examined by polarised light. These appearances are not visible in starch which has been exposed to heat.
- 949. Iodine reaction.—If starch is boiled with water, and a drop of iodine-solution is added to the *cold* liquid, it will assume an intensely *bluish-black* colour.

This colour disappears when the liquid is heated, but it frequently reappears when the liquid is cooled, if the starch is present in large excess.

The colour is destroyed by the addition in excess of a reducing agent, such as H₂SO₃, Na₂S₂O₃, or H₂S: it is therefore prevented from appearing if any one of these substances is present in the starch-solution, until the iodine has been added in excess.

- 950. Br and HCl: if starch is dissolved in a little strong HCl, and Br-water is added, a deep yellow precipitate is formed.
- 951. Basic Lead acetate gives a white precipitate with starch-solution.
- 952. Conversion into Dextrin and Glucose.—When it is boiled with dilute H₂SO₄ for some time, starch is hydrolysed: it is first converted into a mixture of dextrin and glucose, and finally into glucose alone. In the early stages of the process, the dextrin may be detected by giving a red coloration with iodine (954). The glucose may be detected by means of Fehling-solution, after the liquid has been neutralised (908).

DEXTRIN, $(C_6H_{10}O_5)_n$.

953. Dextrin is a white powder, which is soluble in water, but is insoluble in *strong* alcohol. It is therefore precipitated from its aqueous solution by the addition of much alcohol. Its specific rotatory power varies considerably in different specimens.

Dextrin may be made by the hydrolysing starch (952), or by heating starch to 200° for some time. It is a mixture of substances which conform to the general formula $(C_6H_{10}O_5)_n$. It appears to contain the CO group, since it gives a yellow coloration with 'alkalis and can unite with phenylhydrazine.

- 954. Iodine-solution produces a red colour in the solution of most forms of dextrin.
- 955. Fehling-solution (909), and CuA₂ in HA-solution, are usually *slowly* reduced by dextrin-solution on boiling.
- 956. Hydrolysis.—Dextrin is converted into glucose by being heated for some time with dilute H₂SO₄; when this solution is neutralised and heated with Fehling-solution, red Cu₂O is precipitated (908).
- 957. Basic Lead acetate gives no precipitate with dextrinsolution. (Difference from starch.)

GROUP VII.—GLUCOSIDES.

958. Glucosides are compounds of vegetable origin. When they are hydrolysed, they usually yield glucose, $C_6H_{12}O_6$, and an aromatic compound: thus *salicin* gives glucose and saligenin; digitalin gives glucose, digitalose and digitaligenin; and amygdalin splits into glucose, hydrocyanic acid and benzaldehyde.

SALICIN, C13H18O7.

- 959. Salicin occurs in willow-bark; it crystallises in silky needles, and melts at 201°. It has a bitter taste, and is sparingly soluble in cold water and in alcohol, but is more soluble when the liquid is heated. Salicin is insoluble in ether, but is readily soluble in caustic alkalis and in glacial acetic acid.
- 960. Ignition: when salicin is heated in an ignition-tube, it melts, then chars and emits the smell of burnt sugar.
- 961. Strong H₂SO₄, when it is added in small quantity to solid salicin, gives a blood-red coloration.
- 962. Ammoniacal AgNO₃ (Note, 759) and a few drops of KOH-solution, when they are warmed with solution of salicin, give a silver mirror.

963. Hydrolysis: when salicin is heated with dilute H_2SO_4 . it is hydrolysed and yields glucose and saligenin, or ortho-hydroxybenzyl alcohol: $C_{13}H_{18}O_7 + H_2O = C_6H_{12}O_6 + C_6H_4CH_2OH.OH$.

A portion of the hydrolysed liquid is rendered alkaline with KOH-solution and is then warmed with Fehling-solution (909); reduction takes place with separation of orange-red cuprous oxide.

A second portion of the hydrolysed liquid is treated with a few drops of K₂Cr₂O₇-solution, the suspended saligenin is coloured pink.

Powdered K₂Cr₂O₇ and strong H₂SO₄ are added to a third portion of the hydrolysed liquid, the fragrant smell of salicylic aldehyde will be perceived.

If FeCl₃-solution is added to a fourth portion of the hydrolysed liquid, a violet coloration will be produced.

- 964. Mandelin's reagent (1046) produces a purple-red coloration.
 - 965. Froehde's reagent (1045) gives a violet coloration.
- 966. Erdmann's reagent (1044) gives a bright red coloration with a purple edge.

DIGITALIN, C29H46O12.

967. Digitalin occurs in the foxglove. It is a white powder which melts at 217°. Digitalin is slightly soluble in cold water, readily soluble in alcohol and almost insoluble in ether and in chloroform.

By hydrolysis digitalin yields glucose, digitalose and digitaligenin: — $C_{29}H_{46}O_{12} + H_2O = C_6H_{12}O_6 + C_7H_{14}O_5 + C_{16}H_{22}O_2$.

- 968. Strong H₂SO₄ gives a yellow coloration, which gradually changes to brown and finally to red.
- 969. Strong H_2SO_4 and $K_2Cr_2O_7$: if digitalin is dissolved in strong H_2SO_4 and a trace of powdered $K_2Cr_2O_7$ is added, a brown coloration is produced which gradually changes to green.
- 970. Strong H₂SO₄ and Br-water: on adding strong H₂SO₄ and subsequently stirring with a glass rod which has been dipped in bromine-water, a reddish-brown coloration is produced.
- 971. Ammoniacal AgNO₃ (Note, 759) and a drop of KOH-solution produce a silver mirror when the liquid is warmed.
- 972. Mandelin's reagent (1046) produces a brown coloration changing to red.
 - 973. Froehde's reagent (1045) and Erdmann's reagent (1044); each of these produces a brown coloration.

AMYGDALIN, C₂₀H₂₇O₁₁N.

- 974. Amygdalin occurs in bitter almonds and in other vegetables; it has a melting-point of 215°. In contact with water, amygdalin is hydrolysed by an enzyme, "emulsin," which is present in the bitter almond; the hydrolysis yields glucose, hydrocyanic acid and benzaldehyde:— $C_{20}H_{27}O_{11}N + 2H_2O = 2C_6H_{12}O_6 + HCN + C_6N_5COH$.
- 975. Nitrogen: the presence of nitrogen in amygdalin may be proved by the sodium test (679), or by heating it with sodalime (678).
- 976. Hydrolysis: when it is heated with dilute H₂SO₄, amygdalin is hydrolysed, yielding glucose, HCN and benzaldehyde.

The presence of *glucose* in the hydrolysed liquid may be shown by Fehling-solution (909); HCN may be detected by silver nitrate (691), and benzaldehyde may be found by its smell and by Schiff's reaction (875).

GROUP VIII.—ORGANIC BASES.

977. This Section includes the important bases, aniline, toluidine, pyridine and quinoline; also the artificial drugs, acetanilide, phenacetin and antipyrine, and the basic substance urea.

Aniline, C₆H₅.NH₂.

- 978. Aniline is a colourless liquid of peculiar smell, but it usually becomes brown by exposure to the air owing to the presence of an impurity. Aniline boils at 184.4°, has a density of 1.0231, and is slightly soluble in water and in alcohol, and readily soluble in ether and in chloroform.
- · It forms crystalline salts by combination with acids; and these salts are decomposed, with the liberation of the base, by caustic alkalis, but not by AmOH.
- 979. The Salts of Aniline do not give reaction 980; but the base may be liberated from the salts by treating their aqueous solution with KOH, when the aniline will separate in oily drops, and these may be dissolved by shaking the liquid with ether. The

evaporation of this ethereal solution will leave the aniline in the free state, when the test can be applied.

- 980. Strong H₂SO₄ dissolves aniline, yielding a colourless 'solution; if a drop of this solution is added to a little dilute dichromate solution and the solution is stirred, a blue colour will be produced.
- 981. Bleaching-powder, if it is added in small quantity to a very dilute solution of aniline, produces a mauve coloration. If this solution is diluted, and a little dilute Am₂S is added to it, a magenta coloration is produced which is not permanent.
- 982. Chloroform and alcoholic KOH, when they are warmed with aniline, evolve the intensely offensive smell of phenyl isocyanide.
- 983. Bromine-water gives a pale pink precipitate of tribromaniline, $C_6H_2Br_3NH_2$.
- 984. Diazo reaction.—To a cold solution of aniline in dilute HCl five drops of KNO₂-solution are added in a test-tube, the mixture being kept cool by holding the tube in cold water: a few drops of β -naphthol dissolved in NaOH are then added: a brilliant scarlet coloration will be produced, which is due to the formation of benzene-azo-naphthol. This reaction is given by all primary aromatic amines.

TOLUIDINE, $CH_3.C_6H_4.NH_2(1:4)$.

985. Paratoluidine crystallises in lustrous scales which have a faint but characteristic smell. Paratoluidine has a melting-point of 45° and a boiling-point of 200°: it is sparingly soluble in water, but more soluble in alcohol.

Toluidine forms crystalline salts with acids, from which the base can be separated by heating them with caustic alkali solutions.

- 986. Strong H₂SO₄ dissolves toluidine, and the solution gives a yellow colour when a drop is added to dilute dichromate solution.
- 987. HNO₃ gives with toluidine a blue colour, which gradually turns violet, then red, and finally brown.
- 988. Bromine, dissolved in strong HCl, gives with toluidine solution a precipitate of dibromotoluidine.
- 989. Bleaching-powder gives no coloration with toluidine. (Difference from aniline.)

- 990. $\mathbf{H}_2\mathbf{PtCl}_6$ (*Note*, 420): yellow crystalline precipitate of $(CH_3.C_6H_4NH_2.HCl)_2\mathbf{PtCl}_6$.
- 991. Diazo reaction: if a solution of toluidine in dilute HCl is treated as is described in paragraph 984, it will yield a result similar to that obtained from aniline.

Pyridine, C₅H₅N.

992. Pyridine is obtained from bone-oil. It is a colourless liquid, with a strong characteristic smell, which boils at 115.5° and has a density of 0.9893 at 15° C. It is miscible with water in all proportions, giving a liquid with a feebly alkaline reaction, and is soluble in alcohol and in ether.

Pyridine is a strong base, and combines with acids to form salts, which are decomposed by caustic alkalis with liberation of the base.

- 993. Strong H₂SO₄ forms a white pyridine salt, which dissolves in excess of the acid to a colourless liquid.
- 994. Bromine-water, when it is added in excess to a solution of pyridine in HCl, gives a yellow precipitate, C₅H₅NBr₂.
- 995. $\mathbf{H}_2\mathbf{PtGl}_6$ gives with a solution of pyridine in HCl an orange-yellow precipitate of the platinichloride, $(C_5\mathbf{H}_5\mathbf{N}.\mathbf{HCl})_2\ \mathbf{PtCl}_6$.
- 996. The reagents which produce precipitates with the alkaloids (1036 et seq.) also give precipitates with pyridine.

QUINOLINE, C9H7N.

- 997. Quinoline is obtained from bone-oil and from coal-tar. It is a colourless limpid liquid, with a peculiar aromatic odour, and becomes brown on exposure to the air. It boils at 238°, but volatilises in steam from boiling-water; has a density of 1.0944, is sparingly soluble in water, and more readily soluble in most organic liquids. Quinoline burns with a luminous smoky flame. It is a strong base and yields deliquescent salts by combination with acids.
- 998. Strong H₂SO₄: white crystalline precipitate of the sulphate, soluble in excess of the acid.
- 999. H₂PtCl₆ (*Note*, 420) gives with the HCl-solution a yellow precipitate of the platinichloride, (C₉H₇N.HCl)₂PtCl₆.
 - 1000. K₂Cr₂O₇ gives with acid solutions of quinoline a fine Q.A.

yellow precipitate of the dichromate, $(C_9H_7N)_2H_2Cr_2O_7$, which dissolves when the liquid is warmed. (Difference from pyridine.)

1001. The general reagents for the alkaloids (1036-1042) give precipitates with quinoline.

ACETANILIDE, OR ANTIFEBRIN, C₆H₅.NH.CO.CH₃.

- roo2. Acetanilide forms colourless lustrous crystalline plates, which melt at 113° and boil at 283°. It is slightly soluble in cold water, but readily soluble in hot water, in alcohol, in ether and in chloroform.
- 1003. H₂SO₄ and K₂Cr₂O₇.—If a little acetanilide is dissolved in strong H₂SO₄ and powdered K₂Cr₂O₇ is sprinkled on the surface of the solution, red streaks which quickly turn green will be produced.
- 1004. KOH.—If a little acetanilide is heated with KOH-solution, potassium acetate and aniline are formed :— $C_6H_5.NH.COCH_3 + KOH = C_6H_5.NH_2 + CH_3.COOK$.

The aniline thus produced may be recognised by adding a few drops of CHCl₃ and warming the liquid, when the badly smelling phenyl isocyanide will be evolved.

- 1005. Strong H₂SO₄ and Alcohol, when they are warmed with acetanilide, produce ethyl acetate, which is recognised by its fragrant smell.
- 1006. $HgNO_3$ solution, if it is poured upon a little acetanilide in sufficient quantity to moisten it, gives on evaporation to dryness a green mass, which becomes deep red when it is moistened with strong H_2SO_4 .
- 1007. FeCl₃ solution, when it is boiled with acetanilide, produces a turbity but no red coloration, although the FeCl₃-solution becomes darker in colour. (Distinction from antipyrine and phenacetin.)
- 1008. Mandelin's reagent (1046) produces an orange-red solution, the colour of which changes to red and finally to grey.

Phenacetin, or Para-acetamido-ethoxy-benzene, $C_6H_5O.C_6H_4.NH.COCH_3.$

- 1009. Phenacetin forms colourless odourless crystalline scales, which melt at 135°. It is almost insoluble in cold water, fairly soluble in hot water, and readily soluble in ether and in chloroform.
- 1010. Ignition: when it is heated in an ignition-tube phenace-tin melts and then sublimes.

- 1011. Conversion into salicylic acid.—When phenacetin is heated with twice its volume of zinc-dust until it is charred, salicylic acid is produced, and may be found by boiling the mass with a little water, filtering and adding FeCl₃-solution to the liquid, when a violet coloration will be produced (779).
- 1012. Strong H₂SO₄ and alcohol, when they are warmed with phenacetin, cause the smell of ethylacetate vapour to be evolved.
- 1013. HNO₃ diluted with its own volume of water, when it is warmed with phenacetin, produces an orange-red solution which deposits yellow crystals on cooling. If KOH is then added in excess, a red coloration is produced which becomes deeper in tint when the liquid is boiled.
- 1014. H_2SO_4 and $K_2Cr_2O_7$.—If a few drops of strong H_2SO_4 are added to a little phenacetin in a porcelain dish, and some crushed $K_2Cr_2O_7$ is then sprinkled upon the surface, a green coloration will be produced after a time.
- 1015. FeCl₃ solution gives a red coloration when the liquid is warmed.
- ioi6. Mandelin's reagent (1046) produces a pale blue coloration.

ANTIPYRIN, OR PHENAZONE, C₁₁H₁₂N₂O.

- 1017. Antipyrin forms colourless odourless crystals, which have a melting-point of 114°, are soluble in water and in alcohol, but are only sparingly soluble in ether: the solution has a bitter taste. Antipyrin is neutral in reaction, but it is a basic substance and forms salts by combining with acids.
- 1018. Ignition.—When antipyrin is heated in an ignition-tube it melts and then chars, and gives a brown liquid sublimate.
- 1019. Strong $\mathbf{H}_2\mathbf{SO}_4$ dissolves antipyrin, forming a colourless liquid.
- 1020. Strong HNO₃, when it is heated with antipyrin, gives first a yellow and then a deep red coloration.
- 1021. $\mathbf{Hg}_2(\mathbf{NO}_3)_2$ gives a dirty yellowish-green precipitate, and turns deep red and deposits a red precipitate when the liquid is boiled.
- 1022. KNO₂ and H₂SO₄.—If a fragment of KNO₂ and a few drops of dilute H₂SO₄ are added to solution of antipyrin, a bright

green coloration is produced; and if the solution is strong, green crystals are formed.

- 1023. H₂SO₄ and K₂Cr₂O₇: this test, applied as directed for phenacetin (1014), gives a green coloration.
- 1024. FeCl₃ solution, when it is heated with antipyrin, gives a red coloration.
- 1025. Mandelin's reagent (1046) produces a pale blue colour, which is evanescent.
- 1026. Most of the general reagents for the alkaloids (1036-1042) produce precipitates with antipyrin.

UREA, OR CARBAMIDE, CO(NH₂)₂.

- 1027. Urea is a colourless crystalline substance: it is very soluble in water and in alcohol, but is almost insoluble in ether. Urea melts and begins to decompose at 132° C., evolving NH₃: its density is 1·323.
- 1028. Strong HNO₃, if it is added to solution of urea, causes the separation of the crystalline nitrate: under the microscope the crystals are seen to be delicate rhomboidal scales.
- 1029. Strong H₂C₂O₄ solution produces a white precipitate of urea oxalate; this precipitate consists of needle-shaped crystals.

Since urea oxalate is less soluble in amyl alcohol than in water, the test is made more delicate by dissolving both the urea and the oxalic acid in this solvent instead of dissolving them in water.

- 1030. KOH, when it is heated with urea solution, slowly evolves NH_3 and yields K_2CO_3 ; the carbonate thus produced may be detected by effervescing on the addition of an acid.
 - 1031. Hg₂(NO₃)₂ yields a white precipitate in urea solution.
- 1032. KBrO solution causes a brisk evolution of CO₂ and N gases from urea and from compounds containing urea.

The KBrO-solution must be freshly prepared, by adding Brwater to KOH-solution until the mixture retains a yellow colour after the liquids have been thoroughly mixed.

A similar reaction takes place when KBrO-solution acts upon a urate or upon an ammonium salt.

1033. Biuret test.—If solid urea is heated for a few minutes just above its melting-point, biuret is formed and NH_3 is evolved:— $2CO(NH_2)_2 = NH(CO.NH_2)_2 + NH_3$.

If the residue is then allowed to cool and is extracted with water, and the solution is mixed with several drops of CuSO₄-solution and then with NaOH-solution added drop by drop, a *violet* coloration is produced.

1034. Cyanuric acid: when urea is more strongly heated (1033), cyanuric acid is formed and remains as a white residue:— $3\text{CO}(\text{NH}_2)_2$ = $\text{H}_3\text{C}_3\text{N}_3\text{O}_3$ + 3NH_3 . If the residue is boiled with water, and this solution is treated with a few drops of AmOH- and CuSO₄-solutions successively, a purple precipitate is produced.

GROUP IX.—ALKALOIDS.

ro35. The vegetable alkaloids are basic nitrogenous substances, which are capable of forming additive compounds or salts with acids. They usually occur in plants united with organic acids, such as citric, malic, tannic, and meconic; and they are largely employed in medicine on account of their pronounced physiological action. Most of them are extremely poisonous and possess a bitter taste.

The alkaloids are arranged in the following order. First, quinine, quinidine and cinchonine, which are derived from the cinchona bark; then follow morphine, apomorphine, codeine, narcotine and papaverine, which are contained in opium prepared from the poppy-fruit. Then come strychnine and brucine, which are contained in the Strychnos Nux Vomica and the St. Ignatius bean; caffeine or theine contained in coffee and tea; cocaine in cocaleaves; veratrine; atropine, hyoscyamine and hyoscine in Atropa Belladonna; nicotine in tobacco-leaves, and coniine in Conium Maculatum.

The solutions of the alkaloids are alkaline to test-paper, and the solutions of their salts are either neutral or acid in reaction. The free alkaloids are almost insoluble in water, hence solutions of their salts must be employed for trying the liquid reactions.

GENERAL REAGENTS FOR THE ALKALOIDS.

The following general reagents precipitate most of the alkaloids:-

1036. Na₂CO₃ precipitates the hydrated alkaloid from strong solutions of its salts.

1037. Phospho-molybdic acid (Note, page 230) causes a yellow to brown precipitate in the solution of an alkaloid.

- Note.—Phosphomolybdic acid is prepared as follows: The solution of ammonium molybdate in nitric acid (1410, 14) is precipitated by the careful addition of sodium phosphate solution (1409, 16). The precipitate is well washed, and is then dissolved in Na₂CO₃-solution. This solution is evaporated to dryness and the residue is ignited. Water is then added and warmed with the residue, which is finally dissolved by adding a considerable excess of HNO₃.
- 1038. H₂PtCl₆-solution (*Note*, 420), acidified with HCl, gives with solution of an alkaloid-salt a yellow crystalline precipitate, similar to that obtained with solution of an ammonium salt.
- 1039. Iodine dissolved in KI solution gives a reddish-brown precipitate in the solution of an alkaloid.
- 1040. Ignition.—When an alkaloid is heated on platinum foil, it burns with a bright smoky flame; and when it is heated with soda-lime, it evolves NH₃.
- 1041. Tannic acid gives a white or brownish precipitate in an alkaloid-solution.
- 1042. Picric acid gives a yellow crystalline precipitate in an alkaloid-solution.

SPECIAL REAGENTS FOR THE ALKALOIDS.

The following special reagents are also used in identifying the alkaloids.

1043. The most convenient way of applying the following and similar reagents for colour tests is as follows:—

Place a trace of the alkaloid or of its salt upon a white tile or in a porcelain dish, and add a drop or two of the reagent by means of a thin glass rod; then stir the liquid with the pointed end of the rod: the colour will be produced and will be easily seen upon the white surface.

- 1044. Erdmann's reagent is prepared by mixing 6 drops of strong HNO₃ with 100 c.c. of water; 12 drops of this solution are then mixed with 24 c.c. of strong H₂SO₄ (1043).
- 1045. Froehde's reagent is made by dissolving 1 gramme of ammonium molybdate in 100 grammes of strong H₂SO₄ (1043).
- 1046. Mandelin's reagent is prepared by dissolving half a gramme of vanadium chloride or oxide in 100 c.c. of strong H₂SO₄ (1043).
- HgCl₂ and 50 parts of KI in 940 parts of water.

QUININE, $C_{20}H_{24}N_2O_2.3H_2O$.—Use Quinine sulphate, $C_{20}H_{24}N_2O_2.H_2SO_4.8H_2O$ (1050).

- bark. It forms a white crystalline mass when it is crystallised from its solution in alcohol; and is slightly soluble in cold water, but more readily soluble in hot water, and in AmOH, alcohol, chloroform, ether, petroleum-spirit and benzene: the aqueous solution is alkaline and is strongly basic. Quinine has a meltingpoint of 174°C. when it is anhydrous; the crystals melt at 57° and then lose their water of crystallisation.
- 1050. Fluorescence.—When quinine sulphate is being dissolved in water, a few drops of dilute $\rm H_2SO_4$ should be added; the solution is intensely bitter, and gives a pale blue fluorescence. (Difference from cinchonine.)
- 1051. HCl and Ignition.—When quinine is mixed with HCl and evaporated to dryness, it leave a residue which, when it is ignited, evolves purple vapour similar in appearance to iodine vapour.
- 1052. Alkaline hydroxide or carbonate precipitates hydrated quinine from fairly strong solutions of a quinine-salt; this precipitate disappears when the liquid is shaken with ether.

The corresponding cinchonine precipitate is not dissolved by ether.

- 1053. Br-water, or Cl-water, when it is added in small quantity, does not colour a solution of quinine; but if AmOH is afterwards added, an intense *emerald-green* colour appears, which is due to the formation of thallioquinine.
- 1054. Br- or Cl-water and K_4 FeCy₆.—If the addition of Br- or Cl-water to quinine solution is followed by the addition of K_4 FeCy₆-solution and one or two drops of KOH-solution, a *deep red* tint is produced. This colour quickly changes to dirty brown: it is destroyed by HĀ, but reappears when AmOH is cautiously added.
- 1055. Strong H_2SO_4 dissolves quinine to a colourless solution, which becomes green when a crystal of $K_2Cr_2O_7$ is added.
- 1056. Am₂C₂O₄ gives a white precipitate, the formation of which is hastened by shaking the liquid. (Difference from quinidine.)

1057. Mandelin's reagent (1046) gives no change, but on the further addition of a drop of HNO₃ a violet coloration is produced.

QUINIDINE, 2C₂₀H₂₄N₂O₂.5H₂O.

- 1058. Quinidine, when it is crystallised from alcohol, forms white efflorescent needles which contain two molecules of water. Anhydrous quinidine melts at 171.5°: it is soluble in water, in ether and in alcohol.
- 1059. Strong H₂SO₄, when it is poured upon a particle of quinidine, forms a colourless solution, which turns green when a crystal of K₂Cr₂O₇ is added.
- 1060. $Am_2C_2O_4$ gives no precipitate with quinidine. (Difference from quinine.)
- 1061. Br-water (1053), HCl and ignition (1051), and Mandelin's reagent (1057), give reactions with quinidine similar to those given with quinine.

CINCHONINE, $C_{19}H_{23}N_2O$.

- 1062. Cinchonine crystallises in white lustrous prisms, and melts at 255° to a colourless liquid, which partly sublimes when it is further heated. Cinchonine is sparingly soluble in water, but more soluble in alcohol, in ether and in chloroform. Solutions of cinchonine have an alkaline reaction and a bitter taste.
- 1063. Cl-water, or Br-water, when it is added to a cinchonine-solution, produces no change; but on the further addition of AmOH, a yellowish-white precipitate appears. (Difference from quinine.)
- 1064. K₄FeCy₆ gives a yellow precipitate of cinchonine ferrocyanide, which is readily soluble in excess. The precipitate is soluble in hot water, and crystallises in golden-yellow crystals as the solution cools. This is a characteristic reaction.
- 1065. Strong H_2SO_4 and $K_2Cr_2O_7$ (1055), HCl and ignition (1051), and Mandelin's reagent (1057) give with cinchonine changes similar to those given with quinine.

MORPHINE, C₁₇H₁₉NO₃.—Use Morphine hydrochlorate, C₁₇H₁₉NO₃.HCl.3H₂O.

1066. Morphine meconate is present with a pomorphine, codeine, narcotine and papaverine in poppy-fruit, and in opium prepared

from the fruit. Morphine crystallises in colourless prisms, which contain one molecule of combined water: these become anhydrous at 120°, and melt at 230°C.

Morphine is almost insoluble in water, ether, and alcohol; it is slightly soluble in hot water, but readily soluble in chloroform, benzene, and amyl alcohol, and the solution has an alkaline reaction. The salts of morphine are readily soluble in water and in alcohol.

- 1067. KOH or AmOH yields a white precipitate of the base, which is soluble in excess of KOH, but is only sparingly soluble in excess of AmOH.
- 1068. HNO₃.—The strong acid produces a *yellowish red* colour, which does not change to *violet* on the addition of SnCl₂. (Difference from brucine.)
- ro69. FeCl₃ solution: if the neutral solution is added drop by drop to neutral morphine solution, it produces a *dark blue* colour: this colour disappears when an acid is added.
- ro70. Iodic acid, when it is added to morphine or to its salts, yields free iodine. The separation of iodine is shown by the liquid becoming brown; but the iodine is more certainly and readily detected by adding starch-solution, or by shaking the liquid with CS₂ (555, 2, 3). The brown coloration of the solution becomes more intense on the addition of AmOH.

This test is rendered very delicate and characteristic by moistening the solid substance with a solution of one part of iodic acid in fifteen of water, and then adding a solution of one part of starch in four hundred of water; when very dilute AmOH is poured upon the blue solution thus obtained, a coloured ring is seen at the surface of contact of the two liquids; the ring is blue below and brown above.

This reaction serves to distinguish morphine from other organic substances which contain nitrogen.

- 1071. H₂SO₄ gives no colour when it is added to a solution containing morphine; but when a crystal of K₂Cr₂O₇ is dropped into the acid liquid contained in a white porcelain dish, and the crystal is slowly moved about with a glass rod, an *intense green* colour appears. Quinine gives a similar reaction.
- 1072. H_2SO_4 and $K_2Cr_2O_7$.—If solid morphine, or any of its compounds, is dissolved by heating it with a few drops of strong H_2SO_4 , and a minute quantity of HNO_3 is added to the cold solution, a deep red colour is produced. This colour turns to a mahogany tint when a fragment of $K_2Cr_2O_7$ is dropped into the liquid.

- 1073. Froehde's reagent (1045) gives a purple coloration, which becomes green and finally yellowish-brown.
- 1074. Mandelin's reagent (1046) produces a dull purple coloration which changes to grey.

APOMORPHINE, C₁₇H₁₇NO₂.

- 1075. Apomorphine is a white amorphous substance, melting at 295°. It is slightly soluble in water, but readily soluble in alcohol, ether, benzene and chloroform. Solutions of the free base rapidly become green in colour and finally brown.
- 1076. HNO₃ produces a purple-red coloration, which rapidly changes to brown.
- 1077. FeCl₃ solution gives a purple-red coloration, which becomes dark brown when the liquid is heated.
- 1078. Strong H_2SO_4 and $K_2Cr_2O_7$: when a drop of strong H_2SO_4 is poured upon a trace of apomorphine, and particles of $K_2Cr_2O_7$ are then sprinkled on the surface of the acid, an olivegreen coloration is produced, which changes to brown.
 - 1079. Mandelin's reagent (1046) produces a bluish coloration.
- 1080. Froehde's reagent (1045) gives a deep green coloration, which gradually becomes blue.

CODEINE (Methyl-morphine), C₁₇H₁₇NO(OCH₃)OH.

- 1081. Codeine crystallises from water in prisms, which contain one molecule of water. It is fairly soluble in hot water; readily soluble in alcohol, ether, chloroform, benzene and amyl alcohol; but insoluble in ligroin. Its salts are soluble in water. Codeine is a strong base, with a melting point of 155° C., and an alkaline reaction to test-papers.
 - 1082. Strong H₂SO₄ dissolves codeine to a colourless solution, which becomes blue when it is warmed, and reacts as follows:—
 - (1) FeCl₃ produces in the cold H₂SO₄-solution a blue coloration.
 - (2) If cane-sugar is ground up with the H₂SO₄-solution in a mortar, a purple-red coloration appears after a time.
 - (3) When a drop of strong HNO₃ is added to the H₂SO₄-solution, a deep red coloration is produced.
 - 1083. Formalin: when a little codeine is moistered with

ormalin, and a few drops of strong H₂SO₄ are added, a violet colour produced.

1084. Mandelin's reagent (1046) gives a greenish-blue coloraon.

1085. Froehde's reagent (1045) gives a green coloration, rhich changes to blue and finally to yellow.

NARCOTINE, C22H23NO7.—Use Narcotine, or its sulphate.

1086. Narcotine forms lustrous prisms or needles, which have melting-point of 176°. It is almost insoluble in water, slightly oluble in alcohol and in ether, but readily soluble in chloroform. Varcotine is a feeble base, and its salts have an acid reaction, since hey are partially hydrolysed in an aqueous solution.

1087. H₂SO₄: the strong acid gives a *bluish-violet* coloration, which changes to orange; with some specimens a yellow solution produced at once.

When this liquid is gradually warmed, it becomes first orangeed and then bluish-violet, or purple stripes proceed from the edge f the liquid surface: when the liquid is cooled, the colour changes o cherry-red. If the heating is pushed to the initial evaporation f the acid, an intense reddish-violet colour appears.

- 1088. H₂SO₄ and K₂Cr₂O₇.—If narcotine is subjected to the eaction which has been already described for morphine (1072), n intense red coloration is produced.
- 1089. HNO₃: strong nitric acid produces a yellow coloration in he cold liquid: this changes to red when the liquid is heated, but ades to yellow again as it cools.
- 1090. Erdmann's reagent (1044) produces an orange-yellow olution, which turns pink and then rapidly becomes yellow again.
- 1091. Mandelin's reagent (1046) gives an orange coloration, which rapidly becomes pink.
 - 1092. Froehde's reagent (1045) gives a deep green coloration.

PAPAVERINE, C20H21NO4.

1003. Papaverine is a white crystalline substance which melts it like it is insoluble in water, sparingly soluble in ether, n betwee and in cold alcohol, but readily soluble in chloroform and it alcohol. Its salts are soluble in water.

- 1094. Strong H₂SO₄ dissolves papaverine giving a colourless solution, which becomes violet when it is warmed.
- 1095. K₂Cr₂O₇, when it is added to the acid solution, yields an orange-yellow precipitate of the bichromate.
- 1096. Cl-water and AmOH, if they are added successively, yield a reddish-brown colour, which changes to very dark brown.
 - 1097. Iodine-solutions gives a dark-red coloration.
- 1098. Froehde's reagent (1045) gives a green coloration which changes to blue and finally to cherry-red.
- 1099. Mayer's reagent (1047) gives a white or yellowish-white precipitate.

STRYCHNINE, C₂₁H₂₂N₂O₂.

Use strychnine, or its sulphate, dissolved in water to which a drop of dilute H₂SO₄ has been added.

1100. Strychnine is prepared from Strychnos nux vomica, and from the St. Ignatius bean. It crystallises in colourless prisms which melt at 265°. Strychnine is almost insoluble in water, absolute alcohol, ether and benzene, more soluble in dilute alcohol, and readily soluble in alcohol.

The salts of strychnine are soluble in water and in alcohol. All strychnine solutions possess an intensely bitter taste: one part of strychnine in 700,000 parts of solvent can be recognised by its taste. Both strychnine and its salts are *virulent poisons*.

H₂SO₄ and K₂Cr₂O₇.—Strychnine dissolves in strong H₂SO₄ to a colourless liquid. When a fragment of K₂Cr₂O₇ is added to this solution in a porcelain dish, and is slowly moved about by pushing it with a glass rod, a *bluish-violet* coloration is produced which gradually changes to red or yellow.

The presence of morphine, or of a metallic chloride or nitrate, interferes with this reaction. If any of these substances are present, the strychnine may be precipitated by the addition of solution of $\rm K_3FeCy_6$, or of $\rm K_2CrO_4$; the precipitate is filtered off and slightly washed on the filter, and is then stirred with strong $\rm H_2SO_4$.

- 1102. HNO₃: cold strong nitric acid dissolves strychnine without becoming coloured, but the solution acquires a yellow tint when it is heated.
- 1103. Mandelin's reagent (1046) gives a blue colour, which changes to violet. The addition of AmOH changes the colour to rose-red. (Difference from other alkaloids.)

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Brucine, $C_{23}H_{26}N_2O_4.4H_2O$.

Use Brucine or its sulphate, dissolved in water to which a drop of H₂SO₄ has been added.

- 1105. HNO₃: the strong acid yields an *intensely red* coloration, which gradually changes to yellowish-red and yellow when the liquid is evaporated. After all the HNO₃ has been driven off by evaporation, the addition of SnCl₂ to the hot solution produces an intense *violet* colour. (Difference from morphine.)
- 1106. $\mathbf{H}_2\mathbf{SO}_4$: the strong acid produces a rose-coloured solution, which changes to yellow.
- 1107. $\mathbf{H}_2\mathbf{SO}_4$ and $\mathbf{K}_2\mathbf{Cr}_2\mathbf{O}_7$: if the reaction which has already been described for morphine (1072), is carried out with brucine, the liquid becomes red but quickly changes to yellow.
- 1108. Erdmann's reagent (1044) produces a yellow coloration, which changes to intense red.
- 1109. Froehde's reagent (1045) gives a pale red coloration, which changes to deep reddish-brown.
- 1110. Mandelin's reagent (1046) gives a rose coloration, which rapidly changes to orange.

Caffeine, or Theine, $C_8H_{10}N_4O_2.H_2O$.

- III. Caffeine is present in coffee and in tea, and is a colourless substance forming needle-like crystals, which become anhydrous at 100° C., then melt at 230.5° C., and sublime unchanged at 237° C. Caffeine is slightly soluble in cold water, readily soluble in hot water, and fairly soluble in alcohol, in ether and in chloroform. It is a weak base, its salts being hydrolysed by water.
- 1112. Formation of Murexide.—If strong HNO₃ is mixed with even a trace of caffeine and is then evaporated, a yellow residue remains, which assumes an intense purple colour when AmOH is poured upon it.
- III3. Cl- or Br-water, if it is added to caffeine and the mixture is evaporated, leaves a brown residue: this residue dissolves in AmOH, yielding a violet-red solution.
- 1114. Strong H_2SO_4 dissolves caffeine: when powdered $K_2Cr_2O_7$ is added to the colourless solution, a green coloration gradually appears.
- 1115. KOH: when caffeine is boiled with KOH-solution, the characteristic fish-like smell of trimethylamine is perceived.
- 1116. Mayer's reagent (1047) gives no precipitate with caffeine. (Distinction from other alkaloids.)

COCAINE, or Methyl-benzoyl-ecgonine, C₁₇H₂₁NO₄. Use Cocaine hydrochloride.

- 1117. Cocaine is the active alkaloid which is present in coca leaves. It forms colourless prisms, which melt at 98°, and are sparingly soluble in water, but readily soluble in organic liquids. Cocaine is a strong base and its salts are soluble in water.
- III8. Strong H_2SO_4 does not colour cocaine, but the subsequent addition of a fragment of $K_2Cr_2O_7$ produces a brown coloration.
- 1119. Strong KMnO₄ solution gives a purple precipitate of cocaine permanganate.
- 1120. K₂CrO₄ solution, if it is added to cocaine-solution after a few drops of HCl, produces a yellow precipitate. This is a characteristic reaction.
- 1121. Strong H₂SO₄ and alcohol, when they are added to cocaine, cause a fragrant smell of ethyl benzoate to be emitted when the liquid is warmed.
- 1122. When cocaine is heated with an acid, it undergoes hydrolysis and yields ecgonine, methyl alcohol and benzoic acid.

VERATRINE OR CERAPINE, C32H42NO9.

- 1123. Veratrine is an amorphous white powder, which melts at 205°, and is insoluble in water, but soluble in organic liquids. It possesses no smell, but it causes violent sneezing when it is inhaled each in minute quantity.
- 1124. Strong H₂SO₄ produces a yellow coloration: this colour changes to intense red when the liquid is heated.
- 1125. Sugar and H_2SO_4 : when veratrine is mixed with four times its volume of cane-sugar, and the mixture is moistened with strong H_2SO_4 , the coloration is first yellow, then green and finally indigo-blue.
- 1126. Strong HCl gives no coloration to veratrine, but when the liquid is boiled an intense red coloration is produced.
- 1127. Froehde's reagent (1045) produces a red coloration, which gradually becomes green.

ATROPINE, C₁₇H₂₃NO₃.

1128. Atropine, together with hyoscyamine and hyoscine, is present in the deadly nightshade, Atropa belladonna. It is white

and crystalline, melts at 115°, and is almost insoluble in water, but fairly soluble in ether and benzene, and readily soluble in alcohol and chloroform. It forms salts with acids, is very poisonous, and is employed in ophthalmic surgery.

- 1129. Strong H₂SO₄ dissolves atropine; the colourless solution becomes brown when it is warmed, and emits a fragrant smell.
- 1130. Furning HNO₃, if it is added to atropine and then evaporated to dryness over the water-bath, leaves a residue which, when it is treated with a drop of alcoholic potash, gives a violet coloration changing to red.
- 1131. Ba(OH)₂ solution, if it is added to atropine, and the liquid is then evaporated to dryness, leaves a residue which, when it is heated strongly, emits a smell of hawthorn-blossom.
- 1132. HgCl₂ solution, when it is added to an alcoholic solution of atropine, gives a yellow precipitate.
- 1133. Gold chloride: if atropine is dissolved in dilute HCl, and a 5 per cent. solution of gold chloride is added, a precipitate is formed of the aurichloride, C₁₇H₂₃NO₃HCl.AuCl₃: this precipitate has a melting-point of 136°, and fuses when it is heated with water, forming oily drops. (Distinction from hyoscyamine and hyoscine.)
- 1134. Picric acid gives a yellow precipitate, which has a melting-point of 175-176° C.

HYOSCYAMINE, C₁₇H₂₃NO₃.

- 1135. Hyoscyamine is isomeric with atropine, and is converted into atropine by heat. It is sparingly soluble in water, but is readily soluble in organic liquids. It has a melting-point of 108°.
 - 1136. Strong HNO₃ gives a red coloration.
- 1137. Gold chloride solution (1133) gives a yellow precipitate, which has a melting-point of 165°, and which does not melt when it is heated under water. (Difference from atropine.)
 - 1138. Picric acid forms a picrate, melting at 161°-163°C.

HYOSCINE, or Scopalamine, C17H21NO4.

1139. This alkaloid, formerly thought to be isomeric with atropine, is soluble in water and most organic liquids, and melts at 59°C.

1140. Hyoscine is distinguished from atropine and hyoscyamine by the high melting-point (198°–200°C.) of its aurichloride (1137), which does not fuse under hot water. The picrate has a melting-point of 180°–181°C.

NICOTINE, C₁₀H₁₄N₂.

Use nicotine dissolved in five times its volume of water.

- acids in tobacco-leaves. It is a colourless oily liquid, which boils at 247°, undergoing meanwhile partial decomposition. It rapidly turns brown by exposure to the air, and its tobacco-like odour becomes intensified. It distils unchanged in a current of steam, and is readily soluble in water, ether and alcohol. It forms two series of salts.
- 1142. Strong H₂SO₄ is not coloured by nicotine in the cold, but it becomes brown when it is heated with nicotine.
- 1143. Formaldehyde: if a little nicotine is treated with one drop of formaldehyde and then with two drops of nitric acid, a pink coloration is produced. (Distinction from conine.)
- 1144. Mayer's reagent (1047) gives a reddish precipitate of $\mathrm{C_{10}H_{14}N_{2}.HgI_{4}}$ even with extremely dilute solutions of nicotine: this is a very characteristic reaction.

CONLINE, or Propyl piperidine, C₈H₁₇N.

- 1145. Coniine is obtained from the spotted hemlock, Conium maculatum. It is a colourless liquid with a stupefying odour, and is very poisonous. Coniire boils at 166°, volatilises with steam, and is soluble in water and in organic solvents. It is a strong base and forms neutral salts with acids.
- 1146. Strong H₂SO₄ has no action on coniine when cold, but blackens when it is heated with coniine.
- 1147. Strong $\mathbf{H}_2\mathbf{SO}_4$ and $\mathbf{K}_2\mathbf{Cr}_2\mathbf{O}_7$ quickly produce a green colour.
- 1148. Alcohol, CS₂ and CuSO₄. Add two drops of alcohol, one drop of coniine-solution, then two drops of CS₂ and allow the mixture to stand: then add one drop of dilute CuSO₄-solution: a brown colour will be produced. (Difference from nicotine.)

PART III

SYSTEMATIC ANALYTICAL COURSE AND TABLES

1200. Preliminary Remarks.—The analysis of a substance may be undertaken to prove its complete composition and to prove that no other constituents are present, or it may be desired to ascertain only whether any particular substances are present in it.

The procedure to be adopted in the examination and analysis of a substance depends upon the nature of the substance. For this purpose it is to be generally noted whether the substance is a gas (a), a liquid (b), a non-metallic solid (c), or a metal or alloy (d).

- (a) The method of examining gaseous substances is indicated in par. 1201 by giving a statement of the distinguishing properties of some of the more commonly occurring gases; the scheme for the analysis of a gaseous mixture involves a consideration of the properties of the gases which the mixture may contain.
 - (b) The examination of a liquid is described in pars. 1213 et seq.
- (c) The examination of a non-metallic solid will be found in pars. 1215 et seq.
- (d) The examination of a metallic substance is given in pars. 1299-1305.

Special Tables follow, which provide for the analysis of insoluble substances (1306-1341), of silicates (1312-1315), of substances containing cyanogen (1316-1322), of the rarer inorganic elements (1323, 1324), and of organic compounds (1330-1346).

(2)

1140. Hyoscine is distinguished from atropine and hyoscyamine by the high melting-point (198°–200°C.) of its aurichloride (1137), which does not fuse under hot water. The picrate has a melting-point of 180°–181°C.

NICOTINE, C10H14N2.

Use nicotine dissolved in five times its volume of water.

- at 247°, undergoing meanwhile partial decomposition. It rapidly turns brown by exposure to the air, and its tobacco-like odour becomes intensified. It distils unchanged in a current of steam, and is readily soluble in water, ether and alcohol. It forms two series of salts.
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- 1146. Strong H₂SO₄ has no action on coniine when cold, but blackens when it is heated with coniine.
- 1147. Strong H₂SO₄ and K₂Cr₂O₇ quickly produce a green colour.
- 1148. Alcohol, GS₂ and GuSO₄. Add two drops of alcohol, one drop of coniine-solution, then two drops of CS₂ and allow the mixture to stand: then add one drop of dilute CuSO₄-solution: a brown colour will be produced. (Difference from nicotine.)

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Special Tables follow, which provide for the analysis of insoluble substances (1306-1311), of silicates (1312-1315), of substances containing cyanogen (1316-1322), of the rarer inorganic elements (1323, 1324), and of organic compounds (1330-1346).

SECTION VI

DETECTION OF GASES

1201. Gertain Distinctive Properties of each Gas render its detection possible, when it is alone or when it is mingled with other gases. These distinctive properties of each of the more commonly occurring gases are stated below.

The student may proceed to apply these properties as tests for the detection of gases occurring singly and in a state of mixture. The following facts and suggestions will be of use, when such an examination is to be made. All gases are colourless unless it is otherwise stated.

When a mixture of gases is being examined, it should be remembered that those which act chemically upon one another cannot be simultaneously present. Such mutually exclusive pairs of gases are:—O and NO; HCl and NH₃; SO₂ and H₂S; Cl and H₂S; NH₃ and SO₂; H and Cl in daylight.

In testing for the constituents of a gaseous mixture, the colour, the odour if not dangerous, and the combustibility should be noted. An examination of the properties stated below shows that a brief exposure of a part of the mixture to air would detect NO; shaking another portion with water, followed by the examination of separate portions of this solution, would detect HCl, NH₃, SO₂, H₂S, Cl, CO₂; the application of a flame or spark to another portion of the mixture would detect O, H, N, H₂S, CO, CH₄, C₂H₄ and C₂H₂; and an examination of the products of combustion of a combustible gas will often identify the gas. Special confirmatory tests should be applied, when possible, to support the conclusions drawn from the above general tests.

Some gases are difficult to detect when certain other gases are mingled with them in large proportion. Advantage may frequently be taken of their solubility in water or in solution of caustic alkali to remove these interfering gases and to facilitate the detection of insoluble gases.

- Acetylene, C₂H₂, has an offensive smell as it is ordinarily prepared in an impure condition from calcium carbide: it burns with a smoky flame producing water and carbon dioxide (474), and gives with ammoniacal cuprous chloride solution a red precipitate: it is absorbed by Br-water.
- Ammonia, NH₃, possesses a pungent smell, gives dense white fumes with hydrogen chloride gas (426), dissolves in water, and the aqueous solution turns red litmus blue and turmeric brown: it is absorbed by dilute H₂SO₄.
- Carbon dioxide, CO₂, extinguishes flame, is incombustible, makes limewater milky (474), dissolves slightly in water giving a solution which renders lime-water milky.
- Carbon monoxide, CO, is not soluble in water, burns with a blue flame forming no water, and after the combustion carbon dioxide is found by shaking the product with lime-water (474): it is absorbed by a solution of Cu₂Cl₂ in HCl or in AmOH-solution.
- Carbonyl chloride, COCl₂, is a colourless gas with an unpleasant and pungent smell: it is decomposed by contact with water into CO₂ (474) and HCl (543), and yields urea (1027) when it is heated with NH₃.
- Carbonyl sulphide, COS, is a colourless gas, with a resinous smell faintly resembling that of H_2S : it burns with a blue flame which deposits S on a cold surface, and forms CO_2 and SO_2 . This gas is slightly soluble

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- in water, undergoing gradual change into $\rm CO_2$ and $\rm H_2S$: it is quickly absorbed by caustic alkali solution forming carbonate and sulphide.
- Chlorine possesses a peculiar smell and causes coughing, a taper burns in the gas with a red and smoky flame: chlorine dissolves slightly in water, and the aqueous solution bleaches litmus (499, 544); both the gas and its solution turn paper which has been soaked in solution of starch and potassium iodide blue (555).
- Cyanogen, C₂N₂, is an intensely poisonous gas with a disagreeable smell; it burns with a flame of peach-blossom colour, which forms no water: it is somewhat soluble in water, and brownish flocks separate from the solution on standing: it dissolves in alkaline hydroxide solution with production of cyanide and cyanate, which may be identified by tests (689, 697): if H₂S is passed into an aqueous solution of cyanogen it gives a red crystalline precipitate.
- Ethylene, C₂H₄, has a very faint smell, burns with a very luminous and somewhat smoky flame producing water and carbon dioxide (474), decolorises bromine-water, and is slightly soluble in water, but is not absorbed by Cu₂Cl₂-solution.
- Hydrogen burns with a non-luminous flame, which deposits moisture on a cold surface, forming water but no carbon dioxide (474); it is not soluble in water.
- Hydrogen chloride, HCl, fumes in the air, and gives dense white fumes with ammonia gas; it dissolves in water and the aqueous solution reddens blue litmus, and gives a white curdy precipitate when it is mixed with silver nitrate solution (543). HCl is found in presence of Cl by par. 545.
- Hydrogen sulphide, H₂S, possesses a foul smell; it burns with a blue flame which deposits S on a cool surface, and yields water and sulphur dioxide (489); and it dissolves in water, the aqueous solution blackening lead acetate solution (480).
- Methane, CH₄, is free from smell, burns with a slightly luminous flame producing water and carbon dioxide (474), does not decolorise brominewater, and is not soluble in water.
- Nitrogen is incombustible, extinguishes a flame, and gives negative results with all the tests which are mentioned for other gases.
- Nitrogen monoxide, N₂O, is colourless, is unchanged by contact with air, has a sweetish taste, does not colour nitric oxide, kindles a glowing splinter into flame and produces a pale yellowish-green halo around the flame.
- Nitric oxide, NO, is colourless, but becomes reddish-brown when it is mixed with air or with oxygen.
- Oxygen kindles a glowing splinter into flame, and gives a reddish-brown gas with nitric oxide; it is not appreciably soluble in water, but is absorbed by alkaline solution of pyrogallol.
- Ozone, O₃, has a smell resembling chlorine diluted with air. It is a powerful oxidiser, bleaching moist litmus and indigo, and liberating I from KI-solution. Ozone may be distinguished from H₂O₂ and nitrogen oxides by passing unchanged through KMnO₄-solution, and by giving a violet coloration to test-paper dipped in alcoholic solution of tetramethyl-diamino-diphenyl-methane (631, 632).
- Sulphur dioxide, SO₂, possesses a pungent smell; it dissolves in water and the aqueous solution turns blue litmus red, and changes the red colour of a drop of potassium dichromate solution to green (489).

SECTION VII

ANALYSIS OF LIQUIDS AND SOLIDS

Before commencing work by the General Course (1211 et seq.) the following remarks (1202-1210) should be read.

1202. General Remarks.—It is usually possible to determine by the qualitative analysis of a substance not only the constituents which it contains, but also roughly the relative proportions in which some of these constituents are present. Many of these constituents are precipitated in the course of the analysis, and the volume of each of the precipitates will give an indication of the amount of the constituent which it represents.

This rough estimate is made more exact if a known weight of the substance, say one gramme, is used for the analysis, and the volume of any precipitate obtained from its solution is compared with the volume of the similar precipitate which is obtained from a known weight of the constituent element to be estimated.

In some complex mixtures it is not possible to determine in this way the relative amounts of the different components, but a general idea may often be gained by observing the colour, the solubility and other properties of the complex mixture.

The usual procedure in examining a liquid or solid substance may be divided into three parts:—

- 1. The Preliminary Examinations for Metals and Acid-radicles.
- 2. The Wet Examination for Metals.
- 3. The Wet Examination for Acid-radicles.

The detailed directions for these Examinations are tabulated as far as is possible, the requisite precautions to be observed being also stated.

Until the student is familiar with the general procedure it will be found advisable to read completely through each part of the text before commencing the Examination.

PRELIMINARY EXAMINATION OF A COMPLEX SUBSTANCE.

1203. Before the student proceeds to apply the whole of the Systematic Course to the analysis of substances of complex composition, he will do well to examine a few substances for Metals and Acid-radicles by the Preliminary tests only, which are given in paragraphs 1213–1228. For a detailed account of any test in these Tables the student must refer back to the description of that test which is given in Part II.

A little study of the Tables 1218-1228 will show that they contain certain principal tests marked thus—"Exp. I." These stand in the first column. In the second column are placed in succession the results which may be noticed on trying these experiments; a glance through these will forewarn the student of what has to be looked for.

The main classification of the observations which may be made is indicated by black-type letters; the subsidiary divisions are marked by small numbers and italics; and the individual results are simply placed in succession in a vertical column.

Occasionally an observation is made which it is advisable to confirm by an additional experiment. This Confirmatory-test is entered in the first column just below the original observation, and is marked *Confirmatory* in order to distinguish it from the main experiment; its results are placed beside it in the second column.

The student should understand that he may obtain any of the results which are entered in the second column.

FULL EXAMINATION OF SUBSTANCES IN SOLUTION.

1204. After the student has become familiar with the Preliminary Examination of solid and liquid substances, he should fully analyse some easy substances which contain only a few metals and acid-radicles. A beginning is made at paragraph 1215, and after the Preliminary Examination has been finished, the substance is dissolved, and a Systematic Examination of its solution is made.

The following paragraphs (1205-1210) give a general explanation of the method of making and examining the solution for metals, and also state some points which require attention during the procedure which is stated in the General Table (1235).

• 1205. It is necessary first to separate the Metals into Groups from the solution, by adding the Group-Reagents (82) in such order that each reagent precipitates the metals of one Group only, and leaves in solution the metals of all other Groups which may be present.

This is effected by adding the Group-Reagents to the solution of the substance in the order which is directed in the General Table (1235): the metal-groups will thus be precipitated in succession, the metals of the Potassium or Fifth Group alone being left in solution.

The different Metal-groups, in the order of their analytical separation from one another, are those into which the metals have already been divided for the description of their reactions in Part II.

Each of these Groups has been designated by its Group-number, or by its Group-precipitant, or by some well-known member of

the Group. As has been stated (82), the designation by number is the least safe, since some analysts number the groups differently.

The Analytical Groups are as follows and, omitting the rarer metals, they contain the metals indicated:—

Group I., or HCl-group, or Silver-group (Ag, Pb, Hg).

Group II.A.-B., or H₂S-group:—

A. Copper-group (Cu, Hg, Pb, Bi, Cd).

B. Arsenic-group (As, Sb, Sn).

Group III.A., or AmOH-group, or Iron-group (Fe, Al, Cr). Group III.B., or Am_2S -group, or Zinc-group (Zn, Mn, Ni, Co). Group IV., or Am_2CO_3 -group, or Barium-group (Ba, Sr, Ca). Group V., or Potassium-group (K, Na, Li, NH₄, Mg).

Taking these Groups as they stand in the General Table (1235):—

The first Reagent added is HCl, which precipitates only the metals of Group I., and leaves the metals of Groups II., III., IV. and V. in solution. Hence if a precipitate is obtained when HCl is added in excess, it is filtered off; any member or members of Group I. will then be present upon the filter, while the members of all other Groups will remain in the filtrate.

When H₂S is passed into the Filtrate (Note) metals of Group II. alone will be precipitated and may be filtered off.

AmCl and excess of AmOH are added to the Filtrate, (Note) after the H₂S has been removed from it and Fe has been converted into ferric salt by boiling it with HNO₃. The AmOH will precipitate any members of Group III.A., which may be filtered off, while Group III.B. and Mg will remain dissolved by the AmCl, and Groups IV. and V. will also remain in solution.

The addition of Am_2S to the Filtrate (Note) will then precipitate Group III.B. as sulphides, which are removed by filtration.

Group IV. is then precipitated from the Filtrate (Note) by the last Group-Reagent, Am₂CO₃, and any precipitate is filtered off.

The Filtrate, or Solution if Am₂CO₃ has caused no precipitate, is examined for Group V., the members of which are not precipitated by any Group-Reagent, and will therefore now remain in solution if they were present in the original liquid.

Note.—Or into the solution, supposing no precipitate to have been obtained, and filtration therefore to have been unnecessary.

1206. Addition of Reagents in Excess.—Each Metal-Group must be completely precipitated before the next Group-Reagent is added. If any precipitate has been formed by a Group-Reagent, it is accordingly necessary to add that reagent in excess before proceeding to add the next one. Unless the presence of each reagent

in excess is proved before proceeding, much confusion may be caused.

The general method for ascertaining that a Group-Reagent is present in excess consists in adding a few drops more of it to the well-mixed liquid, the liquid being rendered clear for this purpose either by filtration or by letting the precipitate settle. If any further precipitate is formed, the whole of the liquid must be well stirred with more of the reagent, and the clear filtrate must then be again tested in the same way. This process is repeated until no further precipitate is caused by the addition of the Group-reagent.

In certain cases this procedure is rendered unnecessary, since the presence of an excess of the reagent is detectable by its smell, after the solution and the reagent have been well mixed, and the air above the liquid has been blown out. This is the case with H₂S and with AmOH. The presence of an excess of a coloured reagent may usually be seen by the colour of the filtrate from the well-mixed liquid, as in the case of yellow Am₂S. If, however, an acid or an alkaline substance is directed to be added in excess, the student should always prove the presence of an excess by the use of test-papers on the well-mixed liquid.

1207. Washing Group-precipitates.—Before a Group-precipitate is examined, it is necessary that it should be washed until it is free from adhering solution (45-47). This must also be done in the separations by filtration which are described in the Group-Tables. If the washing has not been satisfactorily carried out, many complications may arise.

1208. Evaporation before Precipitating Group III.—Certain organic substances hinder or prevent the precipitation of the metals of Group III.A. by AmOH. These substances will usually have been already detected in the Preliminary Examination. If they have been detected, it is necessary to destroy them by evaporation and ignition before proceeding to precipitate Group III.

Evaporation to dryness and gentle ignition of the residue are also necessary in order to separate SiO₂ (646), since if this remained in solution it might be mistaken for Al(OH)₃. The process also decomposes borates and fluorides.

But if organic substances, borates, fluorides, and silicates are known to be absent, this evaporation is unnecessary, and it is sufficient to boil the filtrate from Group II. until it has no smell of H_2S ; a few drops of strong HNO₃ or of H_2O_2 are then added, and the boiling is continued for several minutes, in order to completely remove H_2S and to convert ferrous salts into ferric salts.

1209. The Separation of the Sub-groups III.A. and III.B. from one another, by the method which is given in the General,

Table (1235), is seldom complete. For an account of more exact methods, the student is referred to paragraphs 1290 et seq.

The method of examining the precipitate in Group III. may also be influenced by the presence of a phosphate in the solution: the change of procedure, and the causes which give rise to it, are explained in paragraph 1292.

1210. General Remarks.—It will be seen that the full course of analysis is so arranged that it serves not only to detect what is present in a substance, but also to prove that all else is absent. This is usually the object in view when the analysis of an unknown substance is undertaken.

The process of analysis may in some cases be shortened by using separate portions of the solution for each Group and test. The plan which is recommended in the General Table, of working throughout with the whole of the solution, is however preferable; since, as has been already stated (1202), this procedure usually enables the analyst to form an opinion as to the relative quantities of many constituents of the substance, and this is usually important.

Attention may be drawn to the extreme importance of the spectroscope for rapidly discovering certain substances, especially when they are present in minute quantity only (67).

The acid-radicles cannot be satisfactorily separated into Groups by precipitants as has been indicated for the metals: hence those acid-radicles which cannot be detected by the Preliminary Examinations (1218–1228) have to be found by the application of special tests.

An Example of the Entry of the Results of an Analysis is given in paragraph 1347; it will be seen that the form adopted is that of the Analytical Tables.

A list of Substances suitable for Analysis is given in paragraphs 1415-1419.

The student will do well at first to gain experience in the analysis of easy mixtures, which are completely soluble in HCl and contain no cyanogen, organic salts, silicates, borates, fluorides, or phosphates precipitable in the Third Group. The analysis of such substances will be simple, since it will be unnecessary to evaporate the filtrate from the Second Group to dryness, or to follow any special method of precipitating and examining Group III.

When the ordinary course of analysis has been mastered, the analysis of substances, in which the above complications exist, should be attempted: and finally minerals, water residues, and artificial products and by products should be analysed, in which minute traces of their constituents should be carefully looked for.

GENERAL COURSE OF ANALYSIS FOR SOLID AND LIQUID SUBSTANCES

The analytical procedure may be influenced by the presence in a substance of cyanogen or of a silicate. Unless these are known to be absent, it is therefore well to make the tests described in paragraphs 1211, 1212 on small portions of the substance.

1211. Cyanogen Test.—A small portion of the substance is subjected to the tests for cyanogen, known as the "Prussian-blue" and the thiocyanate tests and described in paragraphs 692, 694. If cyanogen is detected refer to paragraphs 1316 et seq., for a description of the preparation of the solution and of the method of analysis.

If cyanogen is not present, the substance is examined by paragraphs 1213 and 1214 if it is a liquid, and by paragraphs 1215 et seq., if it is a solid.

1212. Silicate Test.—Silicate is tested for by fusing some of the finely-powdered substance in a bead of microcosmic salt (648). If a liquid is being examined, a portion is evaporated to dryness, and the residue is fused into the bead. If silicate is present, and solution of the substance by acids is found impossible (1229), it may be necessary to refer to paragraph 1312.

I. THE SUBSTANCE FOR ANALYSIS IS A LIQUID.

- 1213. Reaction to Test-paper.—Test the liquid with blue litmus-paper and with turmeric-paper (52): one of the following results will be obtained:—
- 1. It is Neutral, not changing the colour of either paper; this shows the absence of acids and alkalis, and of salts with acid or alkaline reaction. The salts of Ag and Mg and certain salts of Am, Na, K, Li, Ba, Ca, and Sr are the only neutral soluble salts.
- 2. It is Acid, turning blue litmus red: this shows the presence of an acid or of a salt with acid reaction.
- 3. It is Alkaline, turning turmeric brown, and indicating the presence of Am, Na, K, Li, Ba, Sr, or Ca as hydroxide, or of a salt with alkaline reaction.
- 1214. Test for Dissolved Solid.—Evaporate a few drops of the liquid upon platinum-foil, thin glass, or porcelain, smelling it occasionally:—
- 1. A Residue is left.—Note the colour of the liquid (1216): test a small portion for NH₄ by boiling it with excess of KOH (1218 Confy.), and then evaporate a portion of the solution just to dryness in a porcelain dish, without heating the substance after it is dry, and examine the residue by the Preliminary Tables for Metals and Acid-radicles (1218–1228). Examine the larger part of the solution for Metals by the General Table (1235), paying attention to paragraph 1236: reserve the rest of the solution for the Examination for Acid-radicles by paragraphs 1225 and 1268 et seq.

2. No Residue is left.—The liquid must consist of some volatile substance, probably of water. This is either pure, or it contains certain gases or volatile substances, such as $(NH_4^-)_2CO_3$, CO_2 , NH_3 , HCl, Br, H_2O_2 . These may be detected by their smell, by the action of the liquid on litmus-paper, or by special tests (1201, 624-630).

II. THE SUBSTANCE FOR ANALYSIS IS A SOLID.

1215. Examine the General Properties of the Substance.

If the substance is lustrous and metallic, refer to paragraph 1299 for the method of analysis; if it is non-metallic in appearance, proceed to make a preliminary examination of it as follows.

Note down anything that can be learned about the physical properties of the substance by the use of the senses aided by a lens or microscope, and by a magnetised penknife-blade.

Note especially whether the substance consists of more or less sparkling particles with similar geometrical shape, when it is said to be **crystalline**; or whether the particles are irregular in shape and **amorphous**.

Also test its hardness, by ascertaining whether the substance is readily powdered, or easily scratched with the point of a penknife.

If the particles are sufficiently large, and especially if they are crystalline minerals (1419), ascertain the degree of hardness more carefully, since this will be of use in identifying the substance. Very soft minerals, such as talc and steatite, can be scratched and marked by the thumb-nail: harder substances, such as cryolite or cinnabar, are not affected by the thumb-nail but may be scratched by a bronze coin: still harder minerals, such as apatite and fluorspar, can only be scratched by a penknife: very hard substances, such as quartz, cannot be scratched by a knife, but the knife produces a stain-mark of steel upon their surface.

Try whether the substance is magnetic by immersing the tip of a magnetised blade into the fine powder, and seeing whether its particles are attracted. Fe and certain of its compounds are the most powerfully magnetic bodies known, but Mn, Ni, Co and certain of their compounds are also magnetic.

1216. Note also whether the substance possesses any characteristic smell or colour. The following are some of the more commonly occurring coloured compounds:—

Blue: hydrated cupric salts, and anhydrous cobalt salts.

Green: certain Fē salts are pale green; CuCl₂, salts of Ni, manganates and certain compounds of Cr, are intense green.

Yellow: HgO, PbI₂, As₂S₃, CdS, SnS₂, chromates, Fe···-salts.

Red: HgO, HgI₂, HgS, Pb₃O₄; dichromates are orange-red; permanganates are usually reddish-purple.

Pink: salts of Mn are delicate pink; hydrated salts of Co are reddishpink.

Brown: Fe₂O₃, PbO₂; PbO and CdO are light brown.

Black: CuO, MnO₂, Sb₂S₃, FeS, Fe₃O₄.

White: anhydrous salts of Cu and Fe, and many other substances.

*Colourless: a large number of colourless substances are known; these, as well as many of the faintly coloured bodies, yield white powders when they are crushed.

1217. Reduction to fine powder.—After the above examination has been completed, the solid substance is reduced to the finest powder possible, by rubbing it in the mortar.

An ordinary Wedgwood mortar may be used for powdering substances which are fairly soft; but hard rocks and minerals frequently require to be first crushed in a clean bright steel percustion mortar (Fig. 63) by blows of a hammer, and are afterwards pulverised by trituration only in an agate

mortar.

It must be understood that the subsequent chemical examination is very greatly facilitated if the substance has been carefully reduced to an impalpable powder, which does not feel gritty when it is rubbed beneath the pestle or between the fingers.

A portion of this powder is submitted to the further tests which are described in the following Preliminary Examinations for Metals (1218) and for Acid-radicles (1225), using a small separate portion for each test: the remainder is reserved for the fuller method of examination in solution (1229 et seq.).



Fig. 63.

PRELIMINARY EXAMINATION FOR METALS.

If the substance for analysis is a liquid, part of it may be evaporated to dryness, and subjected to the following tests:—

Experiment.	Observation.	Inference.	
218. Exp. I.—Heat a little of the substance in a small test-tube or ignition-tube.	A. The substance does not change. The substances described under B. below are probably absent, or present only in very small quantity.	Absence of organic substances which blacken and give off a smell of burning; of volatile substances which sublime; of substances which are altered by ignition; and of vater combined with or absorbed by the substance, which would be evolved as steam and form drops on the side of the tube.	
į	B. The substance changes. 1. It changes colour: Yellow, hot. white, cold. Yellow, brown , yellow ,	ZnO. PbO. SnO ₂ or Bi ₂ O ₃ . Fe ₂ O ₃ . Presence of organic matter. An acetate. A tartrate, etc. Probable presence of Co-, Mn-, or Cusalts.	
	2. It juses, and becomes solid again on cooling.	Alkali-salts or certain alkaline earth salts.	
Confirmatory. — The	3. It sublimes. White, crystalline, sparkling subli-	Compounds of Am, As, Hg; or free S or I. HgCl ₂ , As ₄ O ₆ .	
sublimate is carefully examined, if necessary with the aid of a lens.	mate. White, non-crystalline sublimate. Black sublimate, becoming red when rubbed.	Probably Am-salt. HgS.	
	Yellow sublimate. Sublimate of reddish drops, \ which are yellow when cold. \ Violet vapour, cooling to blackish	As ₂ S ₃ . HgI ₂ , becomes sear- let when rubbed. S free, or from certain polysulphides. Presence of I.	
Confirmatory. — Heat another portion of the substance, mixed with about three times as much Na ₂ CO ₃ in fine dry powder and a little KCN,	crystals. Grey mirror, which, when rubbed with a splinter of wood or a glass rod, or when examined by a lens, is seen to consist of globules of Hg. Blackish-brown shining mirror, and smell of garlic, no globules.	Presence of Hg.	
In an ignition-tube (140). Confirmatory.— Pour strong KOH-solution upon the substance and heat to boiling; or mix the substance with soda-lime in a mortar, and moisten and	NH ₃ is given off, known by its smell and by turning moist red litmus- paper blue or turmeric-paper- brown.	Presence of NH4.	
heat.	4. The substance gives off water. It fuses first, then gives off water, and again becomes solid if the	Presence of water, absorbed or combined. Water of crystallisa-	
•	heating is continued. It swells up considerably while it is giving off water.	borax and certain borates, alums, and phosphates.	

Experiment.	Observation.	Inference.
Confirmatory. — Examine the drops of water on the sides of the tube with blue and red litmuspapers.	The water is alkaline. The water is acid.	Probably NH ₄ -compounds. Presence of salts of the volatile acids, e.g. HNO ₃ , HCl, H ₂ SO ₄ .
Confirmatory. — Introduce a burning splinter of wood into the tube.	5. Gas or vapour is given off.* a. It is without smell. It burns more brightly, and if introduced with a spark at the end is inflamed.	O [from chlorate, nitrate, peroxide, etc.; or N ₂ O.
Confirmatory. — Introduce a glass rod, moistened with limewater, into the tube.	The flame is extinguished. The lime-water turns milky. The lime-water does not turn milky, but the flame is extinguished.	CO ₂ or N. CO ₂ from carbonate, oxalate, etc. N from AmNO ₂ probably.
Confirmatory. — Introduce a glass rod or slip of paper moistened with $K_2 Cr_2 O_7$ -solution.	b. The gas or vapour has a smell. Smell of burning S. The $K_2Cr_2O_7$ turns green; moist blue litmus is also reddened by the gas.	SO ₂ from combustion of free S, or from acid sulphites, thiosul- phates, reduction of sulphates or oxida- tion of sulphides.
Confirmatory. — In- troduce a glass rod car-	Reddish-brown nitrous fumes are given off, known by their pecu- liar smell, and by not colouring starch paste orange red. Smell resembling Cl:— Yellowish gas, which bleaches moist litmus.	Nitrates of heavy metals, e.g. of Ph, Bi, Hg. (C) from certain chlor- ides.
rying a piece of moist itemus. Confirmatory. — Introduce a glass rod with moist starch-powder on its end.	Brown vapour, which colours the moist starch-powder orange red. Violet vapour, which colours starch-solution blue (552, Note 1). NH ₂ is smelt.	Br from certain bromides, I, free or from certain iodides. NH4- or cyanogen-compounds.
	c. The gas can be ignited, and burns at the mouth of the tube with a:— Pale yellowish green flame, more or less explosive. Bright white flame, producing white fumes; the unburnt gas smells of garlic. Peach-blossom coloured flame.	NH ₃ probably from strongly heated 'NH ₄ NO ₃ . PH ₃ , probably from a phosphite or hypophosphite. C ₂ N ₂ from a cyanide, probably from HgCy ₂ .
1219. Exp. II.—Dip a moistened loop of platinum wire, which gives no colour to the Bunsen-flame, into the substance, and hold the loop with the adhering powder near the top of the Bunsen-flame, or in the inner blowpipe-flame. Then moisten with a drop of strong HCl and heat again. Continue the heating until no further change occurs in the colour of the flame.	A. The substance colours the flame:— 1. Intense yellow. Examine the flame through the indigo-prism; a crimson colour is seen. 2. Pale lilae, crimson through the indigo-prism. 3. Yellowish green. 4. Crimson, same through the indigo-prism.	Na. Presence of K, Sr, Li. The spectroscope is decisive (65). K. Ba probably. Sr, Li.

^{*} This will usually be shown by effervescence of the substance; if the gas or vapour is not detected by its colour or smell, it can only be found on testing for it specially by the confirmatory tests below.

Experiment.	Observation.	Inference.
The flame coloration should also be carefully examined by means of the spectro-	5. Orange red, dingy green through the indigo-prism.	Ca.
scope (64, 65).	6. Bright green. A blue colour is seen after	Cu or B ₂ O ₃ .
Note. If silicic acid is present, the colorations	moistening with strong HCI.	Cu.
present, the colorations for K and Na usually show only after ignition of the powdered substance with powdered CaSO ₄ .	7. Blue $\left\{ egin{aligned} Intense. \ Livid. \end{aligned} ight.$	$CuCl_2$, $CuBr_2$. As, Sb, Pb, $HgCl_2$, $ZnCl_2$, $SnCl_2$.
the finely-powdered substance, in a small	A. The substance decrepitates or crackles. B. The substance deflagrates, or	NaCl and certain other crystalline salts. Chlorate, nitrate, etc.
cavity scooped in a	causes the charcoal to burn rapidly.	
piece of wood-char- coal, in the blowpipe- flame.	c. The substance fuses easily, and is absorbed by the charcoal, or forms a liquid bead. D. An infusible residue is left on the charcoal: see 1, 2, 3, below:—	Salts of alkalis and certain salts of the alkaline earths.
Confirmatory. — De- tach a portion of the cool	1. The residue is white and very luminous.	Probably BaO, SrO, CaO, MgO, Al ₂ O ₃ , ZnO, or SiO ₂ .
residue from the char- coal, place it upon a piece of red litmus-paper, and moisten with a drop of water.	The paper turns blue, showing the residue to be alkaline.	BaO, SrO, CaO, and possibly MgO.
Confirmatory. — Moisten the residue on the charcoal when cool with several drops of Co(NO ₂).	A blue residue, the colour of which does not disappear with intense heat. A pink residue.	Al ₂ O ₃ , and some phosphates, arsenates, silicates, and borates. MgO.
solution, and heat again strongly in the outer blowpipe-flame.	A green residue.	ZnO, or possibly Sn.
Confirmatory. — Heat	2. The residue is coloured. Colour of bead:	
Confirmatory. — Heat a small quantity of the substance in a clear co- louriess borax bead, first	In outer flame. Green, hot, and \ Colourless, or \	Cu.
in the outer, then in the	blue, cold. Blue, hot and Blue, hot and	Co.
inner blowpipe-flame.	cold. Cold. Cold. Grey or opaque.	Ni.
	Brown-red, hot: \ Olive green, hot \ yellow, cold. \ \ and cold. \}	Fe.
,	Green, hot and Green, hot and cold.	Cr. See following Confirmatory test.
	Reddish purple, Colourless, hot hot and cold.	Mn.
Confirmatory. — The presence of Mn and Cr.	Light-yellow mass on cooling.	Cr.
may be confirmed by fusing the substance with Na ₂ CO ₃ and KNO ₃ or Na ₂ O ₂ , on platinumfoil (Exp. 25, 56).	Blue-green mass on cooling.	Mn. (See 1222.)
Confirmatory. — Heat	3. The residue is coloured, or metallic scales or globules are seen. The substance is reduced to the metallic state (1223):—	Au, Ag, Cu, Pb, Sn, Sb, Bi; [Co, Ni, Fe, Mn,
stance, mixed or covered with powdered KCN and Na ₂ CO ₃ or K ₂ C ₂ C on charcoal in the inner	a. Without incrustation.	form grey powders), Au, Ag, Cu, Co, Ni, Fe, Mn, Sn. Ag, Sn: Ni if hard.
charcoal in the miner blowpipe flame.	Brilliant white metal. Yellow metal.	Ag, Sn: Ni if hard. Au.

Experiment.	Observation.		Inference.
•	netised knife-b	racted by a mag- lade (1223).	Cu. Fe, Co, Ni, Mn.
	b. With incrustati	on.	
Confirmatory. — Detach several of the globules with the point of	Globules. White, malleable, not marking paper.	Incrustation. None, or very slight.	Sn.
a knife, and strike them on the bottom of an in-	White, brittle.	Orange, hot.	Bi.
verted mortar with the pestle; if they flatten to a cake they are malle-	White, malleable, marking paper.	Yellow, cold. S Yellow, hot and cold.	Pb.
able, if crushed to a powder they are brittle. Try also if a globule, fixed on the point of a	White, brittle, giving white fumes.	White, close to substance.	Sb.
knife, marks paper.	None.	Yellow, hot.	Zn.
Note.—The appearance and malleability of one	None.	White, cold. Red-brown, easily	Cd.
reduced metal is frequently much altered by the pre- sence of another.	None.	wolatilised. White, and white fumes; smell of garlic.	As.
1221. Exp. 1V.—Fuse some of the substance, mixed with Na ₂ CO ₃ .	coin is found to be stained black (481). Also if a drop of HCl is placed upon the mass, a smell of H ₂ S is perceived and lead-paper is blackened (480).		Presence of S, free or combined.
on charcoal in the inner blowpipe-flame produced from a spirit-lamp, or from a gasflame known to be free from S: remove the fused mass when cold, place it on a bright silver coin, moisten with a drop of water, and let stand for several minutes.			Note. — Since this reaction serves to detect S and sulphuracid-radicles, it more properly belongs to the Preliminary Examination for acid-radicles: it is, however, best tried here.

Notes to the preceding Preliminary Table.

1222. The green colour produced by Mn conceals the yellow colour due to Cr; on boiling the residue with water it gives a pink or purple solution, the colour of which is best seen on filtering: this colour is due to the formation of $\rm KMnO_4$, and shows Mn to be present. On acidifying this solution with HÅ and boiling again for several minutes, the purple colour is destroyed; and on filtering, the yellow colour due to Cr is seen: the presence of Cr may be further confirmed by the formation of a yellow precipitate in the acid solution on addition of $\rm Pb\tilde{A}_2$ -solution.

1223. The metal is best separated and examined by detaching the mass when cold from the charcoal, and crushing it in a mortar or on a watch-glass with a little water, letting stand for a short time, and then quickly pouring off the water down a glass rod or pestle which is wetted and pressed against its edge; by several times repeat. this operation, the heavier metallic particles alone are left in the mortar or watch-glass (61). The particles of metal may be dissolved at once in dilute HCl or HNO₃

and special tests may be applied to the solution for the identification of the metal.

1224. General Note.—When the substance to be examined contains several bodies, they frequently more or less mask one another's reactions: thus Co if mixed with Fe will give a borax bead which is green whilst how and blue when cold, thus resembling Cu, but differing in remaining blue in the inner flame.

Hence the composition of many complex mixtures is only roughly indicated by the Preliminary Examination, and must be confirmed and established in the wet way, or in solution. Many substances, more particularly minerals, however, can be completely analysed by a careful Preliminary Examination.

PRELIMINARY EXAMINATION FOR ACID-RADICLES.

1225. The substance is required for this examination in the state of powder or of strong solution.

•If the substance given for analysis is a solution, part of it may be evaporated to dryness: the dry residue is then finely powdered and subjected to the following tests. If several acid-radicles and metals are present, the reactions may be more or less perfectly concealed or altered, hence failure in obtaining a certain reaction does not in all cases necessarily prove the absence of the corresponding acid-radicle.

Experiment.	Observation.	Inference.
1226. Exp. I.—Treat some of the substance with dilute HCl and note the result, then heat moderately.	One or more of the following gases may be evolved:— A colourless inflammable gas, free from smell. A colourless gas without smell, which turns milky a drop of lime-water on the end of a glass-rod (474). A gas of suffocating smell, which turns a drop of K ₂ Cr ₂ O ₇ solution green (489).	Hydrogen or methane from a metal or metal carbide. CO ₂ from a carbonate. SO ₂ from a sulphite, or from a sulphate in the presence of As ₄ O ₆ or other reducing
	Yellow S separates, and SO ₂ is evolved (494). A gas with fetid odour, which blackens a drop of PbA ₂ solution, or a piece of lead- paper (480).	substance. See 1288. SO ₂ and S from a thiosulphate. See 1288. H ₂ S from a sulphide; or possibly from a sulphite or thiosulphate, if Zn or other reducing agent is present. See 1288.
	Reddish fumes. A yellowish-green gas of suffocating smell, which blearles moist litmus-paper, is evolved (499):— a. In the cold; and also evolved on adding HĀ. b. Only when heated.	From a nitrite. Cl from hypochlorite. Cl from action of HCl on oxidising substances, e.g. MnO ₂ , chromate, uitrate, chlorate, etc.
	A colourless gas, smelling of bitter almonds.	HCN from a cyanide.
Dip into the above HCI- solution a strip of turmeric-paper.	On drying this paper at a gentle heat, it becomes reddish- brown (634).	Presence of a borate.
1227. Exp. II. — Heat another portion of the substance gently with strong H ₂ 804. No attention need be paid to the gases evolved in Exp. I. (1226).	1. A gas is evolved with pungent smell, which fumes in the air, and renders milky a drop of AgNO ₃ -solution made acid with HNO ₃ -	HCl, HBr, HI, HF * from chloride, bromide, icdide, or fluoride. Bromide al o evolves brown Br-vapour, and iodide pur- ple I-vapour (see below).
Confirmatory. — Mix some of the substance with MnO ₂ free from chloride (547), add strong H ₂ SO ₄ and warm.	A yellow gas is given off with strong smell, which bleaches moist litmus-paper. Brown vapour is given off, which colours a little moist starch-powder orange-red. Violet vapour is given off, which colours a drop of starch-paste blue (554).	Probably CI from chloride. Br from bromide.

^{*} HF acts upon the glass, and the SiF4 formed renders the drop milky with SiO2, but the glass

Experiment.	Observation.	Inference,
Confirmatory. — Drop into the hot liquid a few small pieces of copper. If no brown fumes are seen proceed to the next	2. A heavy fuming suffocating gas is evolved, and the liquid behaves as if the glass were greasy: on rinsing out the tube, and drying it thoroughly, the inside is seen to be corroded and dimmed. 3. The acid fumes are reddish, usually seen only on heating after adding a few fragments of Cu.	HF from fluoride: confirm by 654. The HF acting upon the silies of the glass evolves SiF ₂ , which is detected by holding a moistened glass read in the gas; SiO ₂ will be deposited as a gelatinous film upon it (655). Presence of nitrate. If iodide is present it must usually be separated by CuSO ₄ (553) before a nitrate can be detected.
test. Confirmatory.— Boil some of the solid substance with water, or take some of the liquid; cool, add about twice as much strong H ₂ SO ₄ , cool, and pour cold solution of FeSO ₄ carefully in upon the top of the acid liquid.	A dark brown ring or layer forms upon the surface of the acid, either at once or on cooling the liquid by immersing the test-tube in cold water. [Refer to Note, 1276.]	Presence of nitrate.
	4. The acid is coloured reddish- yellow, and a yellow gas is evolved which smells like Ci and bleaches litmus; on heating the acid explosion or crackling occurs. Confirm by 519.	Presence of chlorate. Nots.—For the detection of nitrate or hypochlorite with the chlorate see 524.1287.1286.
	5. The substance changes from yellow to green, O being evolved, as is shown by a glowing taper.	Presence of chromate.
•	 6. Yellowish green gas, with suffocating smell, which bleaches moist litmus. 7. The same gas as in 6, but 	Cl from a hypochlorite (de- tected already in 1226), or from a chloride in pre- scare of MnOg, etc. A chloride in presence of a
	coloured reddish brown by nitrous fumes: or coloured reddish brown by CrOCl ₂ .	nitrate or nitrite. A chloride in presence of a chromate.
	8. Substance does not blacken, but evolves CO which burns with a blue flame, and CO ₂ which turns a drop of lime- water milky.	Presence of oxalate. *.
	9. Substance blackens, and SO ₂ is smelt.	Presence of tartrate.
Sonfirmatory — Ada	10. A smell of acetic acid or vinegar is noticed.	HX from soctate.
a little alcohol to the substance and heat with H ₀ SO ₂	Strong fragrant odour.	Ethyl sociate from an acc-

NOTE ON THE PRECEDING TABLE.

1228. The S present in sulphides which are not decomposed by dilute EC (1226), is detected by fusing the powdered substance with two or three times, as much fusion mixture (Na₂CO₃ + K₁CO₃) in a covered crucible, forling with a little water, and placing a drop of the solution upon a bright silver coin. These dark seam will be produced; or by adding excess of an usid to the solution, when H₁S will be evolved. Smaller quantities of S may be decreased by the formation of a black colour or precipitate on adding. The Solution of the

EXAMINATION FOR METALS IN THE WET WAY.

After the Preliminary Examination of the solid substance has been completed, a portion of the substance is dissolved (1229), and the solution is submitted to the systematic examination, which commences at paragraph 1235. This is often termed "The Examination in the Wet Way."

Process of Solution.

1229. The liquids which are used as solvents for solid substances are (a) water, (b) hydrochloric acid, (c) nitric acid, and (d) aqua regia. The majority of substances dissolve either in water or in HCl, a few require HNO₃, but the use of aqua regia is rarely necessary. The method of applying these solvents to the preparation of the solution of a solid substance is stated below.

Introduce a portion of the solid substance, reduced to an impalpable powder (1217), into a flask (Fig. 21B, page 20) or boiling-tube with distilled water, and heat the water to boiling (1232): if the substance dissolves entirely, proceed to 1235.

If the powder does not dissolve, allow the liquid to stand until the undissolved portion has in great part settled, then decent through a filter: the filtrate is Solution I.:—

the filter m	sk: boil with dised above (1233); tle, and decant th	repeat this pro	allow any residue to settle, decant through coss and finally boil the residue with strong
Filtrate is Residue: heat with dilute, then with strong HNO ₃ (1234) as was directed above for HCl, and decant through the filter:—			
+ 12 1	Filtrate is	Residue may be warmed for a short time with a mixture of strong HNO ₃ (1234) with three times as much HCl; if this does not dissolve it, dilute and decant the acid mixture through the filter:—	
Solution II.	Solution III.	Filtrate is Solution IV.	Residue: wash well with water, dry, and examine as is directed for substances insoluble in water and acids (1306 et seq.).
	1	i	1

1230. Solutions I., II., III., IV. need not be examined separately. Add to I. a few drops of dilute HNO, and if no precepitate is pro-

duced (1236) acidify it with HCl; add also some HCl to Solution III. Any precipitate caused by HCl in these solutions is filtered off and examined by Table I. (1250). The Solutions I., II., III., IV. are then mixed together, and any precipitate produced by mixing them is examined by paragraphs 1306 et seq. as an insoluble substance.

It is best to boil down Solutions III. and IV. considerably, in order to get rid of most of the HNO₃ which they contain, before they are mixed with Solutions I. and II.

The mixed Solutions are then examined by 1235.

1231. If a complex solid mixture is given for analysis, and the only object is to obtain a solution as rapidly as possible, the substance may be at once boiled with aqua regia for a few minutes, then diluted and filtered. The undissolved residue is examined as a substance insoluble in water and acids (1036, et seq.). The filtrate is evaporated nearly to dryness, dilute HCl is added, and the solution is examined by the General Table (1235).

with water or with HCl should be noted down. Notice especially whether any gases are given off.

Water may cause the evolution of oxygen from peroxides of the alkalior alkaline earth-metals, of hydrocarbons from metallic carbides, of acetylene from calcium carbide, carbon dioxide from bicarbonates, hydrogen sulphide or hydrogen phosphide from certain metallic sulphides or phosphides.

Many of the gases evolved by treatment with HCl may be at once recognised by their smell; but a closer examination may be neglected, as these gases will have been already obtained in the Preliminary Examination for Acid-radicles (1226).

If the gases are evolved from the aqueous solution when HCl is added, or when the HCl and aqueous solutions are mixed, the corresponding acid-radicles must have been present in salts soluble in water, probably salts of alkali-metals. If the gases are evolved when the residue insoluble in water is treated with HCl, the acid radicles were present in salts insoluble in water.

separate. These usually consist of PbCl₂, rarely of BaCl₂. The crystals should be filtered off and dissolved in a little boiling water. Ba is easily found by the flame-coloration which is given by the solution (406). The formation of a bright yellow precipitate, when a drop of K₂CrO₄ is added to the solution, proves the Presence of Pb: Ba gives only a pale yellow precipitate.

If Pb has been found, and a residue is left after the original substance has been treated with HCl, the residue may consist of PbCl₂:

this substance may be dissolved by boiling it with water, and the use of HNO₃ may be avoided (1234).

1234. HNO₃ seldom requires to be used in dissolving a substance. This acid should be employed, when necessary, only in very small quantity, since it decomposes the group-reagent H_2S with separation of S unless the solution is dilute and cold (1239): the presence of HNO_3 may accordingly delay or prevent the precipitation of Group II.

The solubility of the compounds of the metals in various solvents is given in the Table of Solubilities (1264). A reference to this Table will often be of advantage after the metals and acid-radicles have been detected by the preliminary examinations, and the solubility of the substance has been determined by the preceding directions, as it will serve to indicate what the substance may contain.

GENERAL TABLE FOR SEPARATION OF METALS

1235. If the substance for analysis is a liquid with neutral or alkaline reaction, examined as is directed below.

In this Table, and in the Group-Tables which follow, it has been supposed that be produced when the reagent is added for its detection; and hence no filtration The "original solution" referred to below is the solution of the substance if

Add to the greater part of the "original solution" HCl; if any precipitate is produced, continue to The filtrate or solution, which must give no further precipitate on unless it is already dilute, and saturated with H₂S by passing the The precipitate may contain-PbCl₂-white. AgCl-white. precipitate the metals (1241): it is then filtered : Hg₂Cl₂—white. The precipitate may contain— The solution or filtrate is diluted (See 1238.) precipitated: as soon as H₂S to smell of H₂S (1244), a little HgS-black Insoluble in KOH and in Am₂S. But if an organic substance, Examine the precipitate by Table I. (1250). PbS-black Bi₂S₃—blac CuS—black gently ignited in the dish. When undissolved residue is filtered -black CdS-yellow Add a few drops of this acid solution Note.—When the hot HCl-solution is cooled, PbCl₂ often SnS-brown of Phosphate; if no precipitate SnS₂—yellow Sp₂S₃—range In KOH As₂S₃—yellow and in [Au₂S₃—black] Am₂S. separates in colourless crystals. Add to the rest of the HCI-solution The absence of Pb is not which may form, as quickly as precipitate is produced by AmOH proved, if it is not precipitated in this group, as PbCl₂ is somewhat soluble, and Pb [Note .- Refer to 1294. may therefore be found only in Group II. Examine the precipitate at once by Table II. (1252). The precipitate may contain... Al(OH)₃—almost colourless. Cr(OH)₃—light green. Fe(OH)₈—reddish brown. Note 1.—Any precipitate formed on boiling the filtrate must be added to the above (1244). Examine the precipitate by Table III.A. (1255). Note 2.—Unless the liquid has been sufficiently diluted some of the above metals may be afterwards precipitated by Am2S in Group III., and cause much confusion.

NOTES ON THE

1236. If the liquid under examination is alkaline or neutral in reaction, HNO₃ commenced. A change of colour from green to purple indicates the *Presence of* by the General Table; if a precipitate is produced, more HNO₃ is added, and is examined by the General Table, commencing with the addition of HCl.

A precipitate which is caused and not dissolved by HNO₃ may be finely divided from a thiosulphate; this precipitate may be recognised by its colour, and by not There may also be precipitated H₂SiO₃ (gelatinous, almost colourless), SnO₂ or ShO₃ (grange)

A full examination of this precipitate for all that it may contain may be made it says insoluble residue which remains is then examined as a substantial public.

INTO GROUPS BY GROUP-REAGENTS.

it requires special treatment (1236); if its reaction is acid, it may be at once

all metals are present: if a metal or group of metals is absent, no precipitate will is necessary, the solution being treated as a filtrate.

a solid, or the substance itself if a liquid, to which no reagent has been added.

add HCl as long as it causes any further precipitate, warm and filter :-

addition of several drops of HCl, is boiled down if necessary (1239): it is then diluted with water (1241) gas through the solution (1374) for several minutes; or sufficient H₂S-water is added to dilute and

and H₂S is passed again into it, or more H₂S-water is added, to make sure that Group II. is completely produces no further precipitate the clear solution or filtrate is boiled in a porcelain dish until it ceases strong HNO₃ or Br-water is then added, and the liquid is boiled. borate, fluoride or silicate, may be present the liquid must be evaporated to dryness and the residue the dish is cool, a little strong HCl is poured upon the residue and heated, then water is added: any

oif (x245).

to some $AmHMoO_4$ -solution in a test-tube, and warm gently; a yellow precipitate shows the *Presence* is formed with $AmHMoO_4$, *Phosphate is absent*.

AmCl, heat it to boiling, then add AmOH in slight excess, boil again, and filter off any precipitate possible, keeping the funnel closely covered with a glass plate (1246): if phosphate is present and a refer to Tables (1256); if phosphate is absent proceed as is directed below:—

et seq., for further directions for precipitating and examining Groups III.A. and III.B.]

The precipitate may contain— ZnS—white.	Add to the filtrate or solution, which must be distinctly yellow or brown (1248), Am ₂ CO ₈ , warm gently and filter:—		
MnS—light-pink. CoS—black. NiS—black.	The precipitate (1249) may contain—	The filtrate may contain Mg, K, Na, Li.	
Examine this precipitate at once by Table III.B. (1257).	BaCO ₃ —white. SrCO ₃ —white. CaCO ₃ —white.	Examine by Table V. (1263),	
	Examine the precipitate by Table IV. (1260).		

GENERAL TABLE.

must be added in slight excess, before the examination by the General Table i *Manganate* (609, 610); but if no precipitate forms, the liquid is at once examined the liquid is heated, and if a precipitate still remains, it is filtered off and the filtrat

sulphur, which separates as a *white* powder from a sulphide, and as a *yellow* powde being separable by standing or filtration; it may be disregarded. metastannic acid (1237), Sb₂O₅, AgCl (all three white), SnS₂, As₂S₃ (both yellow)

by boiling the precipitate with aqua regia, and then diluting the liquid and filtering in water and acids (1306); the aqua regia solution is added to the filtrate from is caused by mixing these solutions is examined by Table I. (1250).

1237. Metastannic acid is precipitated from the solution of an alkaline metastannate by the addition of an acid. It forms a white precipitate, which becomes golden-yellow when it is moistened with SnCl₂-solution. It is also recognised by dissolving in the washing-water, and being reprecipitated when the water runs into the acid filtrate.

1238. It must be remembered that strong HCl also precipitates a strong solution of a barium salt. This precipitate, however, is readily dissolved when water is added and the liquid is heated: it is easily distinguished in this way from AgCl and Hg₂Cl₂.

1239. If $\mathrm{HNO_3}$ or aqua regia has been used in dissolving the substance (1229, 1234), or if the solution smells of Cl or $\mathrm{SO_2}$, the liquid should be holled down considerably, until neither of the above gases is smelt when HCl is added and the liquid is again boiled. The solution is then diluted, and $\mathrm{H_2S}$ is passed into it, or $\mathrm{H_2S}$ -water is added, at once, whether the dilution has caused a precipitate or not (1241).

If As has been detected in the Preliminary Examination, H₂SO₃ should be added, until the liquid smells of SO₂ after it has been holled for a short time. The liquid is then heated for some time short of boiling until it no longer smells of SO₂. This treatment reduces arsenic and stannic compounds

to the arsenious and stannous condition.

In the case of As this reduction is to be recommended, since arsenic compounds are not easily precipitated by H_2S . It is also very desirable to reduce stannic compounds, since SnS_2 , unlike SnS, readily runs through the filter, and being yellow in colour may be mistaken for S. It will of course be necessary to try the special tests 184–186 and 201–203 upon the original solution, in order to ascertain in which state As and Sn were originally present. If any precipitate, other than white sulphur, is produced by boiling with H_2SO_3 , refer to paragraph 1240.

1240. While the solution is being boiled with H₂SO₃, some H₂SO₄ is usually formed. This acid may partially or completely precipitate Ph. Ba.

Sr as white sulphates.

 $\mathrm{H_2SO_3}$ may also precipitate An as a fine powder, which causes the cool liquid to appear blue by transmitted and brown by reflected light; this fine precipitate remains suspended but separates as a black powder when the liquid is boiled.

Examine any precipitate which has been formed by H_2SO_3 by Table (1251).

of their oxychlorides. These precipitates may be disregarded, since H₂S readily converts them into sulphides.

1242. H₂S often produces a fine white precipitate of S, which is caused by the presence of certain oxidising substances, such as Cl, HNO₃, HClO₃, H₂CrO₄, HMnO₄, H₂SO₃, FeCl₃ (1239). This precipitate is white and is not separable by standing or by filtration; it may be neglected.

Great care, however, must be taken not to mistake SnS₂ for S; it resembles S in running through filter-paper, but differs by being distinctly *yellow* in colour; it may often be coagulated by shaking or heating the liquid.

*1243. Certain changes may be observed while H₂S is being passed; they should be carefully noted.

Thus while Pb and Hg are being precipitated by H2S, the precipitates often show characteristic colours (103, 134).

Also a solution, which is coloured reddish-yellow by H_2CrO_4 , becomes green; a solution, which is coloured purple by $HMnO_4$, becomes colourless; and a reddish-yellow ferric solution becomes a pale-green ferrous solution.

Each of these changes is attended by separation of sulphur.

1244. If a yellow precipitate forms when the filtrate or solution which has been saturated with H_2S is boiled, the presence of an arsenic or a stannic compound is indicated.

In this case H₂S should be passed into the boiling liquid, or H₂S-water added, as long as it causes any further precipitate, the precipitation being preceded with advantage by reduction with SO₂ (1239). The precipitate is filtered off and added to any precipitate which has been already obtained by H₂S, and is then examined by Table II. (1252).

1245. SiO₂ and other substances may be present in the insoluble residue which is left after evaporation with HCl.

 SiO_2 is distinguished from any other substance by disappearing when it is heated with HF.

Al₂O₃, Fe₂O₃, Cr₂O₃ may remain undissolved by HCl, if they have been rendered difficultly soluble by the ignition to which the residue has been subjected. They can be dissolved by long-continued heating with strong HCl. The solution thus obtained is added to the main solution, which is to be examined for Group III.A. An alternative method consists in fusing the insoluble residue with KHSO₄, dissolving the cool mass in water and examining it by Table III.A. (1255).

 ${
m BaSO_4}$ and ${
m SrSO_4}$ may also remain undissolved. They are detected by heating some of the residue strongly for a short time in the inner blowpipe-flame upon a loop of platinum wire, then moistening the loop with a drop of strong HCl, and holding it in the outer part of the Bunsen-flame. Crimson-red flashes, which appear red through the indigo prism, show the presence of Sr; a yellowish-green flame-coloration shows that Ba is present. These flame-colorations should also be examined by the spectroscope (64, 65).

The following is a less rapid method of detecting Ba and Sr. Dry the inso hable residue; fuse it on platinum-foil with three or four times as much Na₂CO₃; allow it to cool, then boil the mass with water until it is disintegrated; filter, wash the residue well upon the filter, and examine it by Table IV. (1260) for Ba and Sr only.

1246. Mn and Zn may accompany the precipitate of Fe(OH)₃, Al(OH)₃ and Cr(OH)₃ in Group III.A.

The precipitation of Mn is partly prevented by keeping the liquid from exposure to the air, after AmOH has been added in excess.

But the further precaution should be taken of dissolving the precipitate, which has been produced by AmOH, in HCl, and reprecipitating it by AmOH; the liquid is then filtered, and the filtrate is added to the filtrate which was originally separated from the Group III.A. precipitate.

This treatment of the precipitate should be repeated several times, if small quantities of Zn or Mn have to be looked for in the presence of Al or Fe.

It is better still to precipitate Groups III.A. and III.B. together (1290), and to examine the precipitate by Tables III.c., III.D. (1295, 1296). See rules given in paragraph 1294.

1247. The filtrate or solution, after the addition of AmCl and AmOH, may be coloured blue by Ni, brown by Co, or reddish-violet by Cr(OH)₃, which may have been dissolved by the excess of AmOH.



The Cr(OH)₃ alone need be removed before proceeding to examine for the remaining Groups. It is readily separated, as a pale green, flocculent precipitate by boiling the liquid for a short time in a porcelain dish, a little more AmOH being added if the liquid does not smell of NH₃ after it has been boiled for a short time. This precipitate is filtered off, and the filtrate is further examined for Groups III.B., IV., and V. by the General Table (1235).

The coloration of the ammoniacal liquid, which is caused by Ni or Co,

is not removed by boiling the liquid: it is simply noted down.

1248. A greenish colour may be given to the Am₂S filtrate by a trace of FeS remaining in solution: the FeS will usually be separated as a black precipitate by boiling the liquid: the precipitate may be filtered off, dissolved in HCl and tested for Fe^{··}-salt.

· A brown or dark colour in the filtrate from the Group III.B. precipitate shows that Ni is present, the colour being due to the solution of some of the NiS in the excess of yellow Am₂S; but if Groups III.A.-B. have been precipitated together (1290) the coloration of the filtrate may also be due to Cr(OH)₃.

Before proceeding to precipitate Group IV., the dark coloured filtrate should be poured into a porcelain dish and boiled for some time, adding a little water if necessary to prevent evaporation to dryness: when pale-green (Pr(OH)₃)

may separate, leaving a colourless liquid.

If black NiS settles down and leaves a colourless liquid when the lamp is removed for a few minutes, the liquid is passed through a small filter, and the black precipitate is tested for Ni by fusing the portion of the filter-paper, which is stained by the NiS, into a colourless borax-bead. If the bead is violet or brown while hot and yellow when cold after it has been heated in the outer flame, and becomes grey or opaque after it has been heated in the inner flame, Ni is certainly present, and need not be further tested for in Table III.B. (1257).

The colourless filtrate, from which the Cr (OH)₃ or the NiS has been separated by boiling in the dish, is then examined by the General Table for Groups IV.

and V.

1249. Part of the precipitate, which has been produced by Am₂CO₃, often adheres firmly to the inside of the vessel in which it has been produced. If this is the case, the tube should be rinsed out several times with distilled water, and the precipitate dissolved by pouring in some hot HĀ and causing it to run over the inside of the tube. This solution is added to the HĀ-solution of the Am₂CO₃-precipitate, which is subsequently prepared for Table ÎV. (1261).

Since the precipitation with Am₂CO₃ is often somewhat slow, it is advisable to allow the warm solution to stand for a few minutes before filtering or passing on. If only a slight turbidity is caused, it may be removed by filtration and examined for the metals of Group IV., the spectroscope serving

to detect mere traces of the metals in a strong solution.

1250. The precipitate produced by HCl may consist of PbCl₂, AgCl, and Hg ₂Cl₂: it is washed several times on the filter with boiling water, the first washings being kept apart:—

The first portions of the washings may contain PbCl₂, in solution: if this is present in any quantity, it will be deposited in brilliant colourless crystals as the liquid cools.

Test for, or confirm the presence of Pb, by adding K₂CrO₄ to the washings: a yellow precipitate, readily soluble in KOH, shows:—

Presence of Pb.

d Filtrate:

Filtrate: add HNO₃ until liquid is acid: a white precipitate, which unites into curdy particles when the liquid is shaken or boiled, shows:—

Presence of Ag.

Residue on the filter: dissolve in a little aqua regia, add AmOH in excess, and then HCl in excess: warm the solution and drop into it a small bright strip of Cu: a grey deposit which becomes silver-white when it is rubbed proves:—

Residue: the residue is rinsed into a test-tube (50, a.f.) and

boiled with AmOH, and the liquid is then filtered :-

Presence of Hg' (Mercurosum).

[Another method of testing this residue for Hg will be found in 1253.]

Table G.—Examination of the Precipitate formed by boiling the HCl-solution with H₂SO₃. (See par. 1240.)

1251. This precipitate may contain Au, $PbSO_4$, $BaSO_4$, and $SrSO_4$; the last three substances are perfectly white, and if Au is present it is usually seen by colouring the precipitate brown or black; the Au is also usually visible during precipitation by H_2SO_3 , or after the white sulphates have subsided, since it colours the liquid blue by transmitted and reddish by reflected light.

Ba and Sr can be tested for at once as is directed in the last part of par. 1245, by heating a little of the moist precipitate on a loop of platinum-wire in the inner blowpipe-flame, moistening with HCl, and trying the flame-coloration.

A more trustworthy method of procedure is the following:—Rinse the precipitate into a small porcelain dish, using as little water as possible; add a few drops of strong HA, or dissolve in the liquid a small crystal of H₂T, then add AmOH until it is just in excess, and boil for a few minutes; let the liquid stand for a short time, and decant through a filter:—

Solution: add HA in excess, then K₂CrO₄; a yellow precipitate, soluble in KOH, shows:—

Presence of Pb.

Residue: pour upon the residue in the dish a little HCl and a few drops of finO_a, heat to boiling, let stand, and decaut into a porcelain dish:—

Solution: boil until the smell of Clis removed, and add several drops of fresh FeSO asolution; a blue coloration and reddish precipitate show:—

Presence of Au.

Residue: dry by gently heating the dish, fuse with Na₂CO₂ on platinum-folt, and examine the mass for Ba and Sr as is directed in the last part of par. 1245.

1252. The precipitate produced by H2S in the HCl-solution, after it boiled with a small quantity of yellow Am₂S (Note 1), or with much

GROUP II.A.-COPPER GROUP.

1253. The Residue may contain HgS, PbS, Bi₂S₃, CuS, CdS. After the precipitate has been washed and allowed to drain, remove it from the filter into a porcelain dish (50, d), pour upon it strong HNO₃ sufficient to cover it completely, and heat gently as long as any red funes come off, adding more strong HNO₃ if necessary to prevent evaporation to dryness. Boil off marly all the acid, add a little dilute H₂SO₄, let stand for some time, stirring occasionally, and filter:

Residue may contain HgS (black), PbSO₄ (white); but if white, the absence of Hg is not proved.

Remove the residue into a porcelain dish, using as little water as possible; add a little HA, then AmOH in excess, boil and filter :-

Residue: dry on the filter at a scrape the dried residue off the filter (Note 2), and heat it, mixed with dry Na₂CO₃, in a small ignition-tube; globules of Hg are seen on the inside of the tube when it is rubbed or examined by a lens :--

Presence of Hg" (Marcuricum).

[For another medical, see 1250.]

Filtrate: in Add HÄ excess, then K2CrO4, vellow precipitate :-

> Presence of Ib.

Filtrate may contain Bi, Cu, Cd. Add AmOII in excess, boil and filter :---

Precipitate (not easily seen in dark blue solutions): wash dissolve off the filter by pouring upon it a few drops of boiling dilute HC3, and let this solution drop into large quantity of cold distilled water: a milkiness appearing at once or after stirring and letting stand for time, Roma shows :--

Presence of Bi.

Note.—A pre-cipitate produced by AmOII must always be tested as above for Bi, since Pb and Hg might possibly appear here as white precipitates and be mistaken for Bi.

Filtrate may contain Cu and Cd: if blue, it contains Cu (Note 3). Add HCl until the liquid is acid, then add H₂S in excess, filter and wash quickly, keeping the filter covered with a glass plate; rinse the precipitate into a test-tube, pour upon it some dilute H₂SO₄, boil and filter through a covered

filter (Note 4) :- -

Residue: dissolve by pouring a lit-tie boiling dilute HNO₃ upon the filter, add to the solution AmOH in excess, then HA in excess and a few drops of K. FeCy -solu-tion: a chocolate-red precipitate shows :-

Presence of Cu.

Filtrate: add much HgH-water, or add much water and pass HeS for some time : a yellow precipitate shows :-

Presence of ('d.

Note. -If this precipitate dark - coloured. Note 4) CuS filter it off, boil it once more with HaSO4; filter rapidly and test the filtrate by Has for Cd.

Note 1 .- If traces of Cu have to be tested for, Na2S should be substituted for readily soluble in Am2S.

Note 2 If the residue is too small to be removed, the portion of the paper conin proforming this test refer to paragraph 140.

Note 3 Another method of detecting Cd in the presence of Cu, consists in adding as yellow solphide; Cu may then be tested for in the filtrate, which has been holled

in excess, and K4FeCye (158).

Note 4. Unless this precipitate is filtered without delay in a covered funnel. into the filtrate as CuSO4; it then discolours the CdS precipitate, and may be of His-water into the funnel with each fresh addition of wasting water.

Note 5. White S will always be precipitated here, if yellow ammenium sulphide of the white liquid with benzene. The coloured sulphide is seen at the surface A The detection of As, Sb, Sn by the method given in par. 1254 is simple and The 200 and 207; but these methods need not be resorted to unless Au or Pt.

The 200 and 207; but these methods need not be resorted to unless Au or Pt.

The Pt. or it is removed and washed, and the Sb-deposit is dissolved by

Start and mixed with H.S. water, or divided and seturated with H.S. an orangeus, confirming this freeway of St.

Voic 8:—The published of H.S. to the MIL solution in the General Table (1235)

Lation courting out to the application of heir whether As and Sn were originally

is been washed with hot water, is removed from the filter (50, d) and Ω H-solution, and filtered:—

GROUP II.B .-- ARSENIC GROUP.

- The Filtrate may contain As_2S_3 , Sh_2S_3 , ShS_2 , ShS_2 , ShS_2 , PtS_2 , combined with alkaline sulphide. Add to it HCl until it is acid, and pass H_2S unless the liquid smells strongly; the above sulphides, if present, will be reprecipitated; a mere millicures due to precipitation of Sh may be disregarded. (Note 5.) The precipitate is filtered off and examined as is directed below (1254). See Note 6.
- 1254. The Precipitate is allowed to drain for some time upon the filter in the funnel, and is further freed from water by carefully taking the filter out of the funnel, opening it out, and spreading it upon a pile of three or four filter-papers. The precipitate is then re moved to a small porcelain dish (50, d, f) and heated for some time nearly to boiling with strong fuming HCl: the liquid is then cooled and filtered:—

The Residue will consist of As₂S₃, which is almost insoluble in strong HCl.

broths from the filter at a gentle heat, then mix it with three or four times as much powdered KCN and freshly-dried Na₂CO₃, and heat the mixture in a small bulb-tube, removing any drops of water inside the tube by a twisted piece of filter-paper; a black mirror (181):—

Presence of As. (Refer to Note 8.)

Note.—The As_0S_3 may also be dissolved by heating it with a little fuming HNO_3 ; the excess of HNO_3 is then boiled off and As is detected as H_3AsO_4 by adding AmCl, excess of AmOH and $MgSO_4$ (185); or by addition of $AgNO_3$, and then cautiously neutralising with AmOH (184).

The Filtrate may be examined for Sb and Sn by either I. or II. below:—

I. Place a piece of platinum-foil in a porcelain dish and pour the acid filtrate upon it, then drop upon the foil a piece of Zn; H will come off with effervescence, and if either at once or after a few minutes a black stain appears upon the platinum the Presence of Sb is indicated. Confirm as is directed in Note 7.

Remove the platinum-foil and drop in pieces of Zn: as soon as the bubbles of H cease to be given off, Zn still remaining undissolved, remove the pieces of Zn, rubbing and riusing any dark deposit back into the dish (199); let this deposit settle, decant the liquid and heat the solid deposit with strong HCI for several minutes in a test-tube, dilute with a little water, filter if necessary, and add a few drops of HgCl₂-solution: a white or grey precipitate (202) indicates:—Presence of Sn (Note 8).

[Refer to 1304, 1305, if Au and Pt have to be tested for.]

1I. The acid filtrate is poured into a little hydrogen-flask (Fig. 51, page 82), in which H has been coming off briskly for about five minutes, being produced by the action of a little dilute HCl upon some stick Zn. The H is lighted at the jet, and the inside of a small porcelain dish or crucible lid is pressed down upon the flame: a black stain which is not discoved by solution of bleaching-powder shows:—Presence of Sb.

The residue in the flask is tested for Sn, as is directed in the latter part of I. (above).

Am₂S, since CuS is somewhat soluble in Am₂S; but SnS and Au₂S₃ are much more

taining it may be out up and mixed with the Na₂CO₃. For the precautions requisite

KCN-solution to the blue ammoniacal liquid, and passing H₂S; Cd if present falls down considerably with excess of HNO₃, by adding AmOH in excess, then HA

washed quickly, and examined at once, CuS is liable to be oxidised and to pass entirely overlooked if present in small quantity. It is well to pour a few drops

has been used above. Traces of the sulphides are detected by shaking up a part of contact of the benzene with the acid liquid.

rapid. It is excelled in delicacy and trustworthiness by the methods given in paraor traces of As, Sb, Sn have to be tested for.

heating the foil in a test-tube with a little very dilute HNO₃. When this solution red products will form either at once or after the liquid has stood for some

present in the -ous or -ic condition. If this is not certain, tests 184-186 and 201-

TABLE III.A.—IRON GROUP.

1255. The precipitate produced by boiling, after the addition of AmCl and of AmOH in excess, may contain Al(OH)3, Cr(OH)3, Fe(OH)3: the colour of the precipitate will usually show whether it contains any quantity of Fe(OH)3 (reddish-brown), or Cr(OH)8 (pale green), since Al(OH)3 is colourless. The precipitate may be examined by I. or by II., below.

I. Dissolve the precipitate in a little boiling dilute HCl (50, d, e, f), add to the solution pure KOH- or NaOH-solution until the precipitate remains after the liquid has been stirred, then add more KOH or NaOH, boil for some time and filter:-

- 1. Filtrate may contain Al(OH)₃ dissolved in excess of NaOH.

 If it is green, Cr(OH)₃ is also in solution and must be precipitated by fur-ther boiling in a porce-lain dish, and then be removed by filtration.

Add HCl in slight excess then AmOH in slight excess: a colourless gelatinous precipitate shows :-

Presence of Al.

Note.—NaOH and KOH may contain some Al(OH)₃: the analyst mayst affectain that the Al does not come from this source.

2. Precipitate may contain Cr(OH)₃ and Fe(OH)₃.

Dry upon the filter, and carefully avoid charring the paper; when sufficiently dried the substance will have shrunk considerably into small, hard, dark-coloured pieces; detach these from the filter and heat them with a little fused Na₂CO₃ and KNO₃ upon a piece of platinum-foil. Continue the fusion by heating the under surface of the foil in the blowpipe-flame as long as any frothing occurs, then allow to cool; a yellow mass indicates the presence of Cr(Note 1). Boil the platinum-foil in a small porcelain dish with water until the substance is entirely dissolved, or only a dark brown powder (Fe₂O₃) remains undissolved; take out the foil, allow the powder to settle, and decant the liquid through a filter settle, and decant the liquid through a filter ---

 Filtrate will be yellow if Cr is present (Note 2).
 Add HÅ in excess, shown by the colour becoming reddish,

boil for several minutes, then add PbA₂-solution; a yellow precipitate shows:— Presence of Cr (Note 3).

4. Residue in porcelain dish; dissolve by pouring in a little HCl and boiling, add some water, then a few drops of KCNS-solution; a blood-red coloration shows :-

Presence of Fe (Note 4).

II. An alternative method consists in adding Na₂O₂, little by little, to the above HCl-solution of the group-precipitate, until the liquid is strongly alkaline: or NaOH may be first added in excess followed by H2O2 in excess. The liquid is then boiled until effervescence ceases, and is filtered. The Strate will contain Na₂CrO₄ and NaAlO₂. If Cr is present the solution will be wellow, and part of it may be tested for Cr as is described in column 3 above. The Al may be detected in another portion of the filtrate by column I above. If We is present the residue will be brown Fe(OH), treat this as is directed in column 4 above.

Note 1.—Mn is often precipitated with the Fe, and impatts to the fused mass a bluish-green colour. The presence of Mn should be noted here, as it may be entirely precipitated in this Group, and not detected in Group III.B.

Note 2.—The green colour, caused by Ma, conceals the yellow actour due, to the Philip when the cool mass is dissolved in water and the sention is balled with HA, the manishese coloration is destroyed, and the yellow actour becomes orthography affect any brown, manipaids hydroxide has been referred by filtration.

Note it is may have been originally present us a chromic salt of and

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chromate. Ascertain in what state Cr was present, by boiling some of the original substance with Na_2CO_3 -solution and filtering. A yellow filtrate, which, after it has been acidified with $H\bar{A}$, gives a yellow precipitate with $Pb\bar{A}_2$ -solution, shows that Cr was *Present as a Chromate*. If the Cr was present as a chromic salt, it will not colour the filtrate yellow.

Note 4.—Fe is always detected here as a ferric salt, since the solution has been boiled with HNO₃, in the General Table. In order to ascertain in which condition Fe was originally present, some of the substance under analysis is boiled with HCl in a special apparatus (Figs. 40, 41, page 21), the solution is filtered, if necessary, and is then divided into two parts. To one portion freshlymade solution of K₃FeCy₆ (Note, 279) is added, a dark blue precipitate shows the Presence of Fe" (Ferrosum); to another portion KCNS is added, a blood-red colour shows the Presence of Fe" (Ferricum).

TABLE III.P.—PHOSPHATE GROUP.

1256. If (PO₄)"' has been found in the General Table by the molybdate test in a small portion of the filtrate from Group II., and a precipitate has been formed by adding to the rest of the solution AmCl and AmOH, the AmOH must be added in slight excess, and HCl must then be added with constant stirring to the turbid liquid, in quantity just sufficient to redissolve the precipitate.

A little HA is then added and some saturated AmA-solution:—

a. If a precipitate forms, add FeCl₃-solution drop by drop with constant stirring until the liquid remains red in colour.

b. If AmA causes no distinct precipitate, add a little Am₂HPO₄-solution, and then FeCl₃-solution gradually until the liquid is just reddened (*Note*).

Now boil the turbid liquid in a flask or dish for a few minutes, and filter at once:—

 The Precipitate is rejected if FeCl₂ has produced a precipitate after AmA has failed to do so (a). 2. Filtrate should be free from (PO4) and should give no precipitate with AmCl and If this was not the case (b), dry the precipitate on the filter, and heat it with a fused mixture of Na₂OO₃ and KNO₂ on platinum-foil, boil the cool mass with water and AmOH. Proceed to examine this for Groups III.B., IV. and V. by the procedure in the General Table (1735), and begin by adding Am. 8 to filter:-Filtrate, if yellow, contains Cr: acidity a portion with HA and add PbAs-solution; a Residue of FegOs may be rejected. precipitate Group III.B. Test the HCI-solution of yellow precipitate confirms:the original substance for Fe" and Fe" as is directed in Note 4 (1255). Presence of Cr. Add to the rest of the filtrate much AmCl-solution: colourless gelatinous precipitate shows :-Presence of Al.

Note.—An explanation of the principles upon which this Table depends will be found in paragraphs 1292, 1293.

TABLE III.B.—ZINC GROUP.

1257. The precipitate produced by Am₂S in the solution, or in the filtrate from Group III.A., may contain ZnS, MnS, NiS, CoS. The colour of the precipitate will usually show whether it contains any quantity of NiS or CoS which are black, or consists only of MnS (pink), or of ZnS (white).

The presence of Ni will have been indicated in the General Table (1235) by the blue colour of the filtrate which was obtained after boiling with AmOH, and by the dark brown colour of the filtrate which was obtained after boiling with yellow Am₂S. If the black precipitate, which was obtained by boiling the dark brown filtrate from Am₂S (1248), has been proved by the borax-bead to be NiS, the further tests for Ni in the following Table may be omitted.

Remove the precipitate from the filter with cold dilute HCl in the way described in par. 50, c; take out the paper and allow the liquid to stand, occasionally stirring it well. The precipitate will either dissolve leaving only white sulphur, and the milky liquid may be at once examined by 1259; or a black residue will be left, which must be filtered off and examined by Table 1258, the filtrate being examined by Table 1259.

1258. The Residue is black, and may contain NiS, CoS: examine it by I. and II. below:

I. Examination by the Borax bead. Fuse a small quantity of the residue in a clear borax bead in the extreme tip of the outer blowpipe-flame, and note its colour; then fuse the bead for some time in the inner blowpipe-flame, and again observe its colour:—

Blue bead in both flames shows:—

Presence of Co.

Co need not be further tested for, but if any doubt exists as to the presence of Ni the rest of the residue must be examined as below.

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Brown or yellow bead, when cold after fusing it in the outer fame, which becomes grey or opaque in the inner flame, shows Presence of Ni, Absence of Co.

Unless a trace of Co has to be tested for, no further examination need be made.

A bead which is neither brown nor blue, but of some intermediate hue on cooling from the outer flame, indicates the probable presence of both Co and Ni.

Examine the remainder of the black residue as is directed below for Ni and Co, or for Co alone if Ni has been already detected by par. 1248.

III. Further Examination of the Residue.—Rinse the residue from the filter into a porcelain dish, using as little water as possible: pour in a little strong HCl and boil for some time adding occasionally a small crystal of KClO₃, until the black residue is entirely dissolved or only a small quantity of dark sulphur remains; then boil down nearly to dryness [a blue liquid shows Presence of Co], dilute with a little water, liter if necessary into a boiling tube and pour in KCN-solution slowly until any visible precipitate disappears, or until the liquid is alkaline, boil briskly for several minutes, and add much strong NaOCl-solution; allow the liquid to stand at least ten minutes, and filter. [Any black film on the Inside precipitate below]:—

- 26,5 W 340 26 8 1

Trecipitate (black): wash, and confirm the presence of Ni by fusing some of the precipitate, or the paper stained wifth the precipitate, in a borax bead, first in the outer, then in the inner blowpipe-flame: a bead yellow when cold, becoming black in the their same shows:—

Presence of Ni.

Filtrate must be warmed with more NaOCI, or Br-water, and filtered from any additional precipitate which may form, then evaporated to dryness, and strong ENO₃ poured upon the residue as long as any frothing is caused; this is then evaporated just to dryness, the residue dissolved in water, excess of KOH added to it, and any precipitate filtered off, and this, or the paper stained by it, is fused into a borax bead: a blue bead shows.

Presence of Co.

1259-The Solution or Filtrate may contain Zn. Mn. Boil it in a porcelain dish until it ceases to smell of H₂S, then drop in a small crystal of KClO₃, and boil for several minutes. After cooling add pure NaOH-solution, until after stirring the liquid it turns red litmus-paper blue, then add more NaOH-solution, stir well and filter:—

Precipitate is white at first, but rapidly darkens in the air. Heat a portion of it with fused Na₂CO₃ and KNO₃ on platinum-foil; a bluish-green mass is obtained on cooling, showing:—

Presence of Mn.

Note.—Mn may have been originally present either in the basic or acidic form. Its presence in acidic form is indicated by the colour of the original solution, since in the form of manganate it is green, and as permanganate it is purple; and these colours will have disappeared on boiling the HCl -solution for some time, or on adding $\operatorname{H_2S}$.

Filtrate; pass H₂S, a white precipitate forms, often somewhat discoloured, showing:—

Presence of Zn.

Note.—From a dilute solution of Zn(OH)₂ in NaOH the Zn(OH)₂ is precipitated by boiling; hence after adding NaOH, as is directed above, the liquid must not be boiled, else Zn might be precipitated with the Mn(OH)₂ and escape detection. But by diluting and boiling the above filtrate, it is usually possible to cause any Zn(OH)₂ which it holds in solution to precipitate, especially if the alkalinity of the liquid has been somewhat reduced by the cautious addition of HCl in quantity insufficient to make the liquid acid. If this precipitate is separated by pouring the liquid through a double filter, and is then dissolved off the filter and the sides of the boiling tube by a little boiling HA, ZnS may be precipitated perfectly white from this solution by the action of H₂S.

TABLES IV AND IV.A.—BARIUM GROUP.

1260. The precipitate, which has been produced by Am₂CO₃ and may contain BaCO₃, SrCO₃, CaCO₃, is first examined by flame-coloration, and the results thus obtained may afterwards be confirmed by the wet method, as is directed in par. 1261, or by the quicker method in par. 1262.

Examination by Flame-coloration.—Take a small quantity of the precipitate off the filter upon the end of a glass rod, and dissolve it by moving the end of the rod about in several drops of HCl on a watch-glass; dip into this solution a loop of platinum-wire, which imparts no colour to the flame (62, Exp. 31), and hold the loop for some time in the Bunsen-flame.

Ca will impart to the flame a yellowish-red colour, Sr a crimson-red, Ba a yellowish-green. The red colorations yielded by Ca and Sr are distinguished by viewing the flame through the indigoprism; the Ca coloration then appears dingy green, while the Sr-coloration still appears crimson. Ba is usually found by its coloration remaining after the other colorations have disappeared. When all three metals are present together, their colorations are apt to interfere with one another; but if at any time a red colour is visible through the indigo-prism, Sr is certainly present; if a green coloration is visible without using the prism, the presence of Ba is proved. The flame-coloration should also be examined by the spectroscope (63).

r261. Examination in the Wet Way.—Dissolve the rest of the precipitate off the filter by pouring in a little boiling HĀ, and boil the liquid and pour it again through the filter if the precipitate is not entirely dissolved. If the solution is not acid in reaction, mix more HĀ with it, then pour off a small portion (A) of the solution into a test-tube, and put by the larger portion (B). Cool portion

BARIUM GROUP

These lines run across both pages.

I. An immediate precipitate is produced by CaSO₄ in the cold, showing :—

Presence of Ba.

Portion (B) must then be tested for Sr and Ca which may also be present: examine it as is directed below:—

Add K_2CrO_4 -solution until the colour of the liquid is reddish-yellow, heat, and filter through a double filter-paper, pouring the filtrate through the same filter repeatedly, if necessary, until the liquid is quite clear; then add to the liquid, which must be orange-red in colour indicating excess of $H\bar{A}$ and of chromate, AmOH until the colour changes to pale yellow, then add Am_2CO_3 -solution:—

No precipitate forms, showing:

Absence of Sr and Ca.

A precipitate forms, showing Sr, Ca, or both of them, to be present. Add to the liquid $\mathrm{Am_2CO_3}$ in excess, filter, reject the filtrate and dissolve the precipitate off the filter in as little boiling HA as possible; pour off a small part (C) of this solution, reserving the larger portion (D). To (C) add $\mathrm{CaSO_4}$ -solution and boil:—

A precipitate forms:—
Presence of Sr.

Examine portion (D) for Ca as is directed at (D), Column II.

No precipitate forms:—
Absence of Sr.

Examine portion (D) for Ca as is directed at (D) in Column III.

(A) and add to it several drops of CaSO₄-solution; if an immediate precipitate forms, refer to column I. below, or to IV. (3262): if no immediate precipitate forms heat the liquid to boiling, and if a precipitate then forms refer to II. below, or to V. (1262); if no precipitate forms even after several minutes, test portion (B) by III. below, or by VI. (1262).

-TABLE IV.

II. A precipitate is not formed at once on the addition of CaSO₄, but appears on boiling the liquid, showing:—

Absence of Ba and Presence of Sr.

D. Portion (B) is then examined for Ca as directed below:—

Add dilute H₂SO₄, boil, filter, and reject the precipitate; add to the filtrate several drops of H₂SO₄ and boil; if this causes any precipitate, boil and filter, and again test the filtrate by addition of H₂SO₄ and boiling; repeat this process if necessary. To the clear filtrate, which gives no further precipitate on the addition of H₂SO₄ and boiling, add gradually AmOH until after mixing the liquid by thorough stirring or shaking, it turns red litmus-paper blue, then add Am₂C₂O₄-solution and warm gently; a white precipitate, often appearing only after a time, shows:—

Presence of Ca.

III. No precipitate is produced by CaSO₄, even on boiling, showing:—

Absence of Ba and Sr, and Presence of Ca.

D. Confirm the presence of Ca by making portion (B) alkaline with AmOH, and then adding Am₂C₂O₄-solution; a white precipitate forms, showing:—

Presence of Ca.

-TABLE IV.A.

V. In the Solution or Filtrate test for Sr. Ca.
Dilute with water; then add dilute H₂SO₄,
boil and filter, repeating the process until no
further precipitate is produced.

further precipitate is produced.

The Precipitate: moisten with strong HCl, and test for Sr by the flame-coloration, using the indigo-prism, or the spectroscope (411).

The Filtrate is examined by Column VI.

VI. The Solution or Filtrate may contain Ca.

Add AmOH in excess, then add Am₂C₂O₄ and allow to stand: a white precipitate shows the:—

Presence of Ca.

TABLE V.—POTASSIUM GROUP.

1263. The filtrate, which remains after all the Group-reagents have been added in succession to the original solution, may still contain Mg, K, Na, Li and NH₄.

Since, however, NH₄-salts have been added as Group-reagents. NH₄ must be tested for in the original substance by KOH-solution (1218, I, 3).

Proceed to examine for other members of Group V. as is directed below.

Evaporate the filtrate from Group IV. to dryness in a porcelain dish, scrape out the solid residue, and heat it to redness upon a piece of platinum-foil in the Bunsen-flame as long as any white fumes are seen to be given off when the foil is removed for an instant from the flame (Note 1). All NH4-compounds are thus entirely removed.

Dissolve any residue which remains (Notes 2, 3), by boiling the foil in a test-tube with a small quantity of water, to which several drops of dilute HCl have been added, and divide this solution into two parts:-

I. EXAMINATION FOR Mg.

Before testing for Mg in one portion of this solution, it is necessary to remove from it any traces of Ba, Sr, or Ca which it may possibly contain, and which might else be mistaken for Mg.

Add, therefore, several drops of H₂SO₄ to the liquid, boil for a short time and then, whether a precipitate has formed or not, add a little AmCl, then AmOH in excess, Amou, then amon in excess, then several drops of Am₂C₂O₄; if any precipitate forms, filter and add to the filtrate Na₂HPO₄, and if no precipitate forms at once, let the liquid stand for come time, white writer likes. some time: a white crystalline precipitate shows :-

Presence of Mg.

II. EXAMINATION FOR K. Na AND Li.

In the other portion test for K and Na as is directed below, referring to Note 3 if Li may be present. Flame coloration.—Dip into the solution a loop of platinum-wire, which has been proved not to impart any colour to the flame, and hold the loop in the Bunsen-flame; one of the following results will be observed:—

A bright yellow flame coloration, indicating the presence of Na (Note 4).
Examine this coloration through xamine this coloration chrough the indigo-prism; if it appears red. Presence of K is probable (443, and Note 3): if no red colors is visible K is probably absent, or is present only in very minute

A pale lilac coloration, tion, appearing crimson red through the indigo-prism,

Presence of K. and absence of Na.

It is usual to confirm the results of the examination of the fame coloration by pouring the remainder of the solution upon a watch-glass, adding to it several drops of H₂PtCl₈-solution (420, Ncte), (See Note 5), or a little sodium picrate solution, and athring well for some time; the formation of a yellow precipitate shows:—

Presence of K.

See par. 421, and Note 6 below.

Note 1.—If the residue is small in amount, it may be ignited in the dish; but this is not to be generally recommended, since the porcelain dish is liable to be cracked by the heat, and it is also difficult to get entirely rid of the NH salts by heating the residue in percelain

quantity.

Not 2 11 is not safe to place much reliance upon an examination of the

foil for the detection of a small quantity of the transparent residue, but it may usually be detected by producing a crackling noise while the foil is cooling immediately after its removal from the flame. Should there be any doubt, the foil must be boiled with water and a drop of dilute HCl, and the solution examined for Mg, K, and Na, as is directed in the above Table.

If Li is to be separated and detected refer to Note 3.

Note 3.—If Li may be present and is to be tested for, the above residue (Note 2) must be treated in such a way as to convert the metals into chlorides.

It is dissolved in a little water and BaCl₂-solution is added drop by drop as long as it produces any precipitate: then Ba(OH)₂-water is added until the liquid is strongly alkaline to test-paper: the liquid is then boiled and filtered: the filtrate is made just acid with dilute HCl, and the Ba is completely removed by adding AmOH and Am₂CO₃ in excess and filtering. The filtrate is evaporated to dryness, and the residue freed from Am-salts by ignition on Pt-foil. The residue is once more dissolved in a little water, and mixed with Am₂CO₃-solution, and any precipitate of BaCO₃ is filtered off. The clear liquid is once more evaporated to dryness, and the residue is ignited to free it from Am-salts.

The residue, in which only chlorides are present, is now extracted with a mixture of absolute alcohol and ether: the residue left on evaporation of this extract is dissolved in a few drops of dilute HCl, and this solution will

give the red flame-coloration and spectrum-band of Li (443).

The residue left undissolved by the ether and alcohol is then dissolved in a little water and a drop of dilute HCl, and tested for Mg, K and Na. It must be remembered that Li cannot be distinguished from K by examining their flame-coloration through the indigo-prism, but only by the spectroscope (443).

Note 4.—The examination of this flame-coloration by means of the spectroscope (63) should be made if possible.

A yellow flame coloration, more or less intense, will almost always be obtained here, since most substances and reagents contain small quantities of Na. Hence the student must note the intensity of the coloration, and judge from it whether the quantity of Na is small or large. The test for a trace of Na is usually made upon the original substance (1219).

Note 5.—If iodine is present, H₂PtCl₆ will produce an intense red coloration. Hence if iodine is suspected to be present, the above solution should be evaporated to dryness with a little strong HNO₃, and the residue dissolved in a few drops of dilute HCl before it is tested for K with H₂PtCl₆.

This is of course unnecessary, if the solution has already been evaporated to dryness with HNO₃ before precipitating Group III. in the General Table. In case of uncertainty, add H₂PtCl₆ to a drop only of the above solution, and if iodine is found to be present proceed as is directed above.

Note 6.—Mere traces of K and Na may be detected by adding $\mathbf{H}_2\mathrm{PtCl}_6$ and evaporating the liquid to dryness in a porcelain dish upon a waterbath. Absolute alcohol is then poured into the dish and stirred. Any yellow residue shows the *Presence of K*.

The solution will give the pure Na-coloration, if Na is present: and if the yellow residue is filtered off and washed with absolute alcohol, it will

vield the pure lilac flame-coloration of K.

EXAMINATION FOR ACID-RADICLES

1264. In conducting the Examination for Acid-radicles, much time and trouble may usually be saved by considering which of these radicles can possibly be present. Their number may usually be considerably reduced by a reference to the Table of Solubility (1266), since the analyst knows at this stage both the solubility of the substance and also what metals it contains. The number of acid-radicles to be tested for may then usually be further reduced, by a consideration of what radicles have been proved to be present or absent by the preceding Examinations (1267).

1265. The following Table of Solubility only includes the commonly occurring compounds of the more abundant metals. The solubility of other substances may be obtained by reference to special books.

The Table is thus arranged. In a horizontal line at the head are

1266. SOLUBILITY OF THE OXIDES

	Acetate	Arsenate	Arsenito	Borate	Bromide	Carbonate	Chlorate	Chloride	Chromate	Cyanide	Ferricyanide	Ferrocyanide	Fluoride	Hypophosphite
Silver Mercurous Lead Mercuric Bismuth Cuprous Cupric Cadmium Stannous Stannic Arsenious Antimonicus Antimonicus Antimonic Ferric Zinc Manganese Cobalt Nickel Vickel Caleium Strontium	W.3 W. W. W. W.5 W.	D.N. D.N. D.N. D.H. D.H. D.H. D.H. D.H.	D.N. D.N. D.H. D.H. D.H. W. B.H. D.H. D.H. D.H. W. B.H. D.H. D.H. W. W. W. W. W. W.	D.N. D.N. D.H. D.H. D.H. D.H. D.H. D.H.	i. C.N. W.3 W.3 D.H.5 C.H. W. W.5 W.5 W.5 W.5 W.5 W.5 W.5 W.5 W.5	D.N. D.N. D.H. D.H. D.H. D.H. D.H. D.H.	W.3 W. W. W	i. C.N. W.3 W.5	D.N. D.H. D.H. D.H. D.H. D.H. D.H. D.H.	O.N. D.H.5 O.N. D.H.5 O.N. D.H.5 W.3 D.H.6 W.3 W. W. W. W. W.	i. C.N. W.S. W. D.H. i. C.H. C.H. W. W. C.H. D.H. i. w.	1. C.N. D.N. D.H. D.H. i. C.H. C.H. C.H. C.H. C.H. C.H. C.H	W. D.N. D.N. D.N. D.N. D.H. O.H. W.3 W. — W. W. W. J. D.H. I. D.H. L. D.H. U.H. W.	W. W

w = water. n.n. = collect hydrochloric sold. o.n. = concentrated hydrochloric sold.

1 Spice and Sb205 are only slowly attacked by concentrated fich.

2 mae solubility in HCl of the cyline Al203. Cr.O3, Te203 depends on their degree of hydration : after

placed the names of the more commonly occurring acid-radicles and of oxygen in alphabetical order. In the vertical column on the left and right is a list of the metals in the order of the Analytical Groups.

*The Solubility of a Compound is found by following the horizontal line of the name of metal, until the vertical line headed by the acid-radicle is reached: the solubility of the compound will be indicated by the letter or letters standing at this point. The signification of these letters is explained at the foot of the Table where also will be found further information concerning certain compounds in numbered notes. It should be understood that the solubility indicated in the Table is that first arrived at by applying the solvents successively in the order stated in par. 1229, and that this would generally imply solubility also in the succeeding solvents there used. The letter which indicates the solubility of the compound will be found in a horizontal line with the acidradicle portion of the name of the compound.

AND SALTS OF THE METALS.

. Lodide	Nitrate	Nitrite	Oxalate	Oxide	Phosphate	Phosphite	Silicate	Sulphate	Sulphide	Sulphite	Thiocyanate Sulphocyanide	Tartrate	Thiosulphate	
1. C.N. W.3 C.N. Q.H.5 C.H. W. S. D.H. S. W. S. W.	W. W.5 W.5 W.5 W.5 W.5 W.5 W.5 W.5 W.5 W	W.3 W.	D.N. D.N. D.H. D.H. D.H. D.H. D.H. D.H.	D.N. D.N. D.N. C.H. D.H. D.H. D.H. D.H. D.H. D.H. D.H	H.H.H.H.	D.H. D.H. D.H. D.H. D.H. D.H. D.H. D.H.	D.N.8 i. D.H.8 D.H.8 D.H.8 D.H.:i. D.H.8	W.3 D.N. 1. D.N. 1. D.H.5 W.5 W. W. W.5 W. W. W. W.5 W.	D.N.8 D.N. A.R. C.N. C.N. C.H. C.H. C.H. C.H. C.H. D.H. D.H. D.H	D.N. D.N. D.H. C.H. D.H. D.H. D.H. D.H. D.H. D.H	C.N. A.R. C.N. D.H. D.H. C.H. W.3 W.	D.N. D.N. D.N. D.N. D.H. D.H. D.H. D.H.	W.3 W.3 C.H. W. W. W. W. W. W. W. W. W.	Silver Mercurous Lead Mercuric Bismuth Cuprous Cupric Cadmium Stannous Stannic Arsenious Antimonic Antimonic Aluminium Chromic Ferric Zinc Manganese Cobalt Nickel Calcium Strontium Barium Magnesium Magnesium Sodium
w. w.	w. w.	w.	w.	₩.	w. - w.	W.	w.	w.	w. w.	w. w.	w. w.	w. w.	W.	Potassium Ammonium

D.N = dilute nitric acid. C.N. = concentrated nitric acid. A.R. = aqua regia. i = insoluble in acids.

⁶ Anhydrous chromic halides and sulphate are insoluble in acids.
7 These solubilities apply to the freshly precipitated salts. Ferro- and ferricyanides, which are insoluble in acids; are readily decomposed by alkalis.
8 Attificial silicates, which have not been strongly ignited, are decomposed by dilute HCl, gelatinous silies separating if the solution is concentrated. Some natural silicates, which are hydrated, are decomposed by the result of the solution is concentrated. posed by strong HCl, but many silicates are unacted on by acids.

The Way in which this Table is used may be illustrated by an example. In order to ascertain the solubility of zinc sulphate, it is only necessary to glance down the vertical column with sulphate at its head, and on the horizontal line with zinc stands the letter w; this indicates that zinc sulphate is soluble in water, and would therefore have been dissolved for analysis by the first solvent used (1229): but no information is given or is required as to its solubility in the other solvents.

The Application of the Knowledge furnished by the Table may be illustrated by the following example.

In a substance, which was entirely soluble in water, the metals K, Ba, Ag were found. A glance along the horizontal lines opposite the names of these metals shows at once which acid-radicles may be present.

All acid-radicles might be present combined with K, since all its salts are soluble in water. But the presence of Ba, in a substance which is soluble in water, shows that the acid-radicles (SO₄), (PO₄), (CO₃), (BO₃), (AsO₄), (AsO₃), (CrO₄), and (F) cannot be present, since these acid-radicles form compounds with Ba which are insoluble in water. Ag excludes in addition the radicles (S), (Cl), (Br), and (I). Hence among the more commonly occurring acid-radicles, (NO₃) only need be tested for.

This example shows how much the examination for acid-radicles may often be simplified, when the metals which are present in a substance and the solubility of the substance are known.

1267. Several Acid-radicles will also probably have been tested for in the Preliminary Examinations: but in case the results have been interfered with by the presence of other substances, the analyst should refer to the reactions for acid-radicles (460 et seq.), and select decisive tests.

The Examination for Metals may also have yielded proof of the presence of certain acid-radicles. Thus:

A green alkaline solution, becoming purple when it is diluted or acidified (1236), shows

A reddish-yellow solution, becoming green and depositing sulphur when H₂S is passed (1243), shows.

A purple solution, becoming colourless with H₂S (1243), shows

Presence of (MnO₄)"

A yellow precipitate of As₂S₃ with H₂S, which appears only when the liquid is boiled (1244), shows

Presence of (AsO₄)"

An insoluble residue of SiO₂, after the filtrate from the H₂S-group has been evaporated to dryness (1245), shows.

And a vellow precipitate obtained with

Presence of (SiO₃)"

• And a yellow precipitate obtained with AmHMoO₄, before precipitating Group III. (1235), shows

Presence of (PO₄)""

If the presence of any of these acid-radicles has been detected in the above manner, its presence should, if necessary, be confirmed by a characteristic test.

Of the Acid-radicles which remain to be tested for, some can be detected by the plan drawn out in paragraphs 1268-1273, others are most easily found by applying special tests to the original substance (1274 et seq.).

Special Tests for Acid-Radicles in the ${ m Na\,_{\circ}CO_{\,3}}$ -solution.

1268. Removal of Metals. Before the following tests are applied, it is advisable to remove from the substance any metals, other than those of Group V., which it may contain, since some of them might be precipitated by the reagents which are added for the detection of the acid-radicles. If alkali-metals alone are present, this separation is unnecessary, since they are not precipitated by any of the reagents.

The removal of the metals may usually be effected by boiling a portion of the finely-powdered substance with pure Na₂CO₃-solution, which must be added to the clear solution as long as it causes any precipitate. The precipitated carbonates are then removed by filtration, and the clear filtrate is divided into five equal portions.

One of these portions is reserved in case of accident, and a second is kept for the tests for organic acid-radicles. The other three portions are acidified, while they are hot, by the addition of HCl, $\rm HNO_3$ and $\rm H\bar{A}$ respectively; they are then examined as is directed on the next page, a separate part being used for each test.

1269. Certain Metals cannot be Completely Precipitated as Carbonates by boiling the substance with $\rm Na_2CO_3$ -solution: the presence of these metals in the solution is, however, frequently of no consequence. If they must be precipitated, the addition of $\rm Am_2S$, or the passage of $\rm H_2S$, will usually separate them as insoluble sulphides; but the subsequent separation of the excess of $\rm Am_2S$ or $\rm H_2S$, by gently warming the solution, is trouble-some.

But since the analyst knows at this stage of the analysis what metals are present, it should not be difficult to separate them, or to allow for their presence while the tests for acid-radicles are being tried.

1270. The Three clear Acidified Portions of the Filtrate, obtained by boiling the substance with sodium carbonate-solution (1268), are tested by paragraphs 1271-1273. Further special tests (1274-1282) are then tried, if necessary.

Some general tests for organic acid-radicles occurring singly or

together will be found in paragraphs 1343-1346.

1271. Portion I. acidified with HCl. Reagents added:—	Present.
BaCl ₂ -solution, a white precipitate insoluble on boil ing (Note 1) BaCl ₂ -solution, a semi-transparent precipitate insoluble on boiling (Note 2) AmCl and Am ₂ CO ₃ , a semi-transparent precipitate. FeSO ₄ -solution, a dark blue precipitate: FeCl ₃ -solution, a dark blue precipitate: FeCl ₃ -solution, a dark blue precipitate: FeCl ₃ -solution, a red coloration, destroyed by pouring into HgCl ₂ -solution.	$(SO_4)''$ $(SiF_6)''$ $(SiO_3)''$ $(FeCy_6)'''$ $(FeCy_6)^{1v}$ $(CNS)'$

1272. Portion II. acidified with HNO ₃ . Reagents added:—	Present.
AgNO ₃ , a pure white precipitate, easily soluble in AmOH (Note 1)	(Cl)', (BrO ₃)' (526), (IO ₃)' (531)] (Br)' (I)'

1273. Portion III. acidified with $H\overline{A}$. Reagents added:—	Present.
PbA2 solution, a yellow precipitate (Note 5) CaCl2 solution, a white gelatinous precipitate CaCl3, a white pulvarulent precipitate. FeCis solution, a yellowich white precipitate	(CrO ₄)" (F)' (1281) or (FeCy ₆) ^{1v} (1317) (C ₂ O ₄)", probably. (PO ₄)"', or (AsO ₄)"'' (1278, 279).

Note it.—Unless the Na₂CO₅ solution, which was used in preparing the solution for these tests, was tree from 15th, and (CL), these acid racheles, if lettered; may have been present out a saminfration in the Na₂CO₅ and not in the longitude substance: portrops for the common substance should then be septed by paragraphs 1274, 1275.

Note 2.—The presence of (SiFe) should be confirmed by adding KCl to

another part of the solution (468), or by heating the $BaSiF_6$ precipitate or the original substance with strong H_2SO_4 (469).

Note 3.—If AgNO₃ yields a black precipitate, this shows the presence of sulphide, or possibly of thiosulphate. Add HNO₃ and boil; the black Ag₂S will thus be decomposed, leaving a milky liquid, in which any other precipitate is readily seen after the sulphur has been coagulated by heating or by shaking the liquid.

Note 4.—If chloride, bromide and iodide have to be tested for, in the absence of CN (I2II), the Na₂CO₃-solution is examined by pars. 564-566: or the precipitate produced by adding AgNO₃-solution in excess is treated as is directed in par. 563. It must be remembered that (CN), (CNO)' (FeCy₆)^{iv}, (FeCy₆)''' and (CNS) are also precipitated by AgNO₃. Accordingly, if cyanide has been detected (I2II, I27I) the examination for acid-radicles precipitable by AgNO₃ must be conducted by I283-I285.

Note 5.—If a white precipitate of PbSO₄ is produced here, it may be dissolved by adding AmOH in excess and warming the liquid (104); red basic lead chromate will remain if a chromate was present.

SPECIAL TESTS FOR ACID-RADICLES.

1274. Sulphate.—A portion of the original substance is boiled with HCl, and the liquid is decanted or filtered, if necessary, and is tested with BaCl₂; a white precipitate shows the *Presence of Sulphate*.

The presence of sulphide, sulphite and thiosulphate with sulphate may be detected by par. 1288.

1275. Chloride.—A portion of the original substance is warmed with HNO₃, and the solution is decanted or filtered, if necessary, and is tested by AgNO₃; a *perfectly white* precipitate, which is easily soluble in warm AmOH, shows the *Presence of Chloride*.

The presence of other acid-radicles which are precipitable by $AgNO_3$ is detected by pars. 1283-1285.

1276. Nitrate.—The clear aqueous solution or extract of the substance (Note) is mixed with its own volume of strong H₂SO₄, and the liquid is cooled. Freshly-made cold solution of FeSO₄ is poured upon the surface of the acid liquid without mixing; a brown layer or ring at the surface of contact of the liquids (509) shows the Presence of Nitrate.

Note.—If nitrite is present, it may be detected and removed as is directed in par. 525, or it may be removed by boiling the substance with excess of HA before the above test is applied.

Chlorate may be detected in the presence of nitrate by pars. 524 or 1287.

1277. Cyanide, if present, will have been detected by the special

test (1211), and by the smell of bitter almonds, given off by the substance after H₂SO₄ has been added (1226, Exp. 1).

Ascertain in what form cyanogen is present by paragraphs 1284,

1278. Arsenate.—This acid-radicle is unlikely to be present unless As has been detected during the examination for metals, in which the presence of (AsO_4) is rendered probable by the precipitation of yellow As_2S_3 occurring only when the acid liquid is boiled after it has been saturated with H_2S (1244).

If As has been found amongst the metals, proceed to test for (AsO₄) by adding HCl to a portion of the Na₂CO₃-solution (1268) until it is acid, then heating to expel CO₂; and adding AmCl, AmOH in excess, and MgSO₄. Filter off any precipitate which forms when the liquid is warmed and shaken, and pour a few drops of AgNO₃-solution upon the white precipitate on the filter; a change of colour to brown shows the *Presence of Arsenate*.

1279. Phosphate.—Boil some of the original substance with dilute HNO₃, and add a little of the clear solution to some AmHMoO₄-solution: stir the liquid well, and if no precipitate forms, warm very gently; a yellow precipitate shows the Presence of Phosphate.

If (AsO₄) has been detected (1278) the above test for (PO₄) is only trustworthy when the yellow precipitate has been obtained either in the cold or by employing a very gentle heat. If any doubt is felt concerning the presence of (PO₄), boil some of the original substance with H₂SO₃-solution (1239), and examine for PO₄ after the (AsO₄) has been entirely separated by passing H₂S into the boiling HCl-solution.

A precipitate which has been obtained with AmHMoO₄ may be further tested by dissolving it in AmOH, adding AmCl and MgSO₄, filtering, and dropping AgNO₃-solution upon the precipitate; if phosphate is present the white precipitate turns yellow, if arsenate is present the precipitate becomes brown.

1280. Borate.—Warm a portion of the substance with a little dilute HCl, dip into the solution a strip of turmeric-paper, and dry the paper in a steam-oven or at a gentle heat. The change of colour of the turmeric to reddish-brown, which becomes blueblack when the paper is moistened with AmOH, shows the Presence of Borate.

vessel for about ten minutes. Now remove the wax, and if the lines which were traced are etched upon the glass, the *Presence of Fluoride is shown* (654).

If SiO₂ is known to be present (1212), the test for fluoride must be made by conducting the gas, which is evolved when the substance is heated with strong H₂SiO₄, into dilute AmOH; a deposit of gelatinous H₂SiO₄ proves the *Presence of Fluoride* (655).

- 1282. Sulphide.—If the presence of sulphide has been indicated by the tests in pars. 1221, 1226 or 1228, or if the presence of a sulphide is suspected and these tests have not detected it, the substance may be boiled with KOH-solution for some time, and alkaline $Pb\bar{A}_2$ -solution may be added to the clear liquid; the appearance of a black precipitate or coloration indicates the *Presence of Sulphide*.
- 1283. Cyanide, Chloride, Bromide and Iodide.—The cyanide is first tested for by the Prussian-blue test (692) or by the thiocyanate test (693, 694).

If cyanide is found it may be removed by (a) or (b).

- (a) Pass $\rm CO_2$ through the slightly alkaline solution until the escaping gas is free from HCN and no longer gives a turbidity with a solution of $\rm AgNO_3$ containing $\rm HNO_3$.
- (b) Mix the solution with $NaHCO_3$ and heat until the HCN ceases to be evolved in the escaping gas as tested by (a).

Chloride, bromide and iodide may now be tested for in the residual liquid, which will be free from cyanide, by par. 564, 565 or 566.

1284. Cyanide, Ferrocyanide, Ferricyanide and Thiocyanate.—Remove the cyanide as is described above (1283), testing the escaping gas for cyanide as is there directed.

Then acidify the liquid with HCl and add to one portion FeCl₃ in excess: a deep blue precipitate shows the *Presence of Ferrocyan-ide*. Filter off this precipitate: a red filtrate shows the *Presence of Thiocyanate*.

Add FeSO₄-solution to the rest of the acidified liquid: a dark blue precipitate indicates the *Presence of Ferricyanide*.

1285. Chloride, Bromide and Iodide, with Cyanide, and Complex Cyanides: add AgNO₃-solution in excess, filter, and dry and ignite the precipitate; all the cyanogen compounds will now have been decomposed, and the silver halides will remain practically unchanged. Heat the ignited precipitate with melted fusion mixture (563) and test the solution of the cool mass for chloride, bromide and iodide by pars 564-566.

1286. Hypochlorite and Chlorate: test for hypochlorite by adding dilute H₂SO₄, when chlorine gas will be evalved in the cold (499).

Remove the hypochlorite by adding solution of NaOH and then

solution of MnSO₄ (500):--

$$NaOCl + Mn(OH)_2 = MnO_2.H_2O + NaCl :$$

a brown precipitate of hydrated MnO₂ will form, filter this off and precipitate the chloride in solution by adding AgNO₃ in excess.

Filter off the AgCl, and add zinc and dilute H₂SO₄ to reduce the chlorate to chloride; then filter and add more AgNO₃, a white precipitate will indicate the original presence of chlorate.

1287. Chloride, Nitrate and Chlorate: test a portion of the solution for chloride by adding AgNO₃: a white precipitate shows the *Presence of Chloride*.

To the remainder of the solution add Ag₂SO₄-solution in excess to precipitate the chloride as AgCl: filter and boil the filtrate with KOH to expel any ammonium salts which may be present, add Zn-dust and boil again: if nitrate was present it will be reduced, and ammonia will be given off.

Filter off the residue, add HNO₃ in excess and AgNO₃: a white precipitate of AgCl shows that chlorate was originally present

and has been reduced to chloride.

1288. Sulphide, Sulphite, Sulphate and Thiosulphate.— Remove the sulphide by shaking the solution with lead carbonate or with cadmium carbonate: the presence of sulphide will be shown by the lead darkening or the cadmium turning yellow.

Filter, and add BaCl₂ in excess to the filtrate to precipitate the sulphite and sulphate. Filter off the precipitate of these mixed barium salts, reserving the filtrate, and treat the precipitate with ECF to decompose the BaSO₃: a white residue denotes sulphate.

Filter from the BaSO, and add to the filtrate Br-water and BaCl.: a forther white precipitate of BaSO, shows that sulphite was originally present.

The thiosuphate is tested for in the reserved filtrate from the prixed barrium salts by acidifying it with HCl and warming: if wellow sulphur is precipitated and sulphur dioxide is produced, the presence of thiosulphate is shown.

crucibles are used the Teclu or the Méker burner (3) will give a sufficiently high temperature for most fusions.

· Now pour some water into the cool crucible, and either allow it to stand, or boil the liquid, until the solid mass is loosened from the crucible. Boil distilled water in a porcelain dish with this mass, crushing it by pressure with a pestle if it does not quickly fall to pieces.

Allow the residue to settle, pour off the solution through a filter, and boil the residue with a little more water; then pour off the solution through the same filter, adding this filtrate to the former one.

The examination of the undissolved residue is described in paragraph 1309, and that of the filtrate in paragraph 1310.

1309. Insoluble Residue on the Filter.—Wash the residue well with boiling water, then make a hole in the bottom of the filter, and rinse the residue through into a test-tube by pouring upon it a little boiling dilute HCl (Note 1).

The residue will dissolve entirely when it is heated with the acid, if the fusion has been continued sufficiently long (Note 2). Filter, if necessary, and examine the solution by the General Table (1235), bearing in mind that only the metals in paragraph 1306 are likely to be present (Note 3).

Note 1.—If Ag and Pb are known to be present by the Preliminary Examination (1307, II.), dilute HNO_3 must be used instead of HCl for dissolving the residue.

If effervescence is produced by the action of the acid upon the residue, the presence of Ba, Sr, Ca, Mg is indicated.

Note 2.—If the residue does not wholly dissolve, either the fusion was incomplete or the residue was not thoroughly washed. The insoluble residue should therefore be dried again, then finely powdered and treated with fresh fusion-mixture (1308). If it consists of chrome iron ore, it may be dissolved by heating it with HNO₃ and KClO₃.

Note 3.—The Solution should be Evaporated quite to Dryness before Group III. is precipitated, and a water-bath should be used towards the end of the evaporation if the substance spirts: since if SiO₂ was present in the substance, the silicic acid may pass into the solution, and if it is not completely separated here by the process of evaporation (646), its presence may give rise to complications in the analysis.

- 1310. Aqueous Solution of the Fused Mass.—Divide this solution into two parts (a) and (b).
- (a) Add HCl to one part until the liquid is distinctly acid, and evaporate in a porcelain dish to dryness, finishing the process on a water-bath, if necessary, to avoid spirting. Then continue to heat the dish gently over the flame or upon a sand-bath until the residue is quite dry: pour in a little strong HCl and warm, dilute and heat again; an insoluble residue shows:—Presence of SiO₂

Filter off SiO₂ if necessary, and examine the filtrate by the General Table (1235). All may be found here, but other metals, such as Cr, Mn, Zn, Sn, Sb, As, should also be tested for, since their oxides are soluble in alkalis, and may therefore pass into this solution.

(b) Test separate portions of the other part of the aqueous solution for the following acid-radicles as is described below. It must be remembered, however, that unless the fusion-mixture was free from chloride, sulphate and phosphate, the tests for these acid-radicles may be of little value.

Chloride: acidify with HNO₃ and add AgNO₃; white precipitate, easily soluble in AmOH.

Sulphate: acidify with HCl and add BaCl₂; white precipitate insoluble on boiling.

Chromate is indicated by the yellow colour of the solution, and its presence is confirmed by obtaining a yellow precipitate after acidifying with HA and adding PbA₂-solution. The chromium may have been present acting as a metal or as a constituent of the chromate acid-radicle (*Note* 3, 1255)

Phosphate: acidify with HNO₃, add a few drops to some AmHMoO₄-solution, and warm gently; a yellow precipitate forms, often only after a time or when the liquid is gently heated.

If As has been found when H_2S was passed into the hot HCl-solution, the presence of (PO_4) must be confirmed in a portion of the filtrate from which the As has been completely precipitated (1279), else the above yellow precipitate may have been due to (AsO_4) .

Fluoride: add HCl in excess to a part of the solution, stir well, and let stand until the CO₂ has escaped; then add AmOH in excess, then CaCl₂-solution as long as it produces any precipitate, and let stand for a time.

Filter off and dry any precipitate which forms, and pour strong H₂SO₄ upon it in a platinum or lead crucible covered with a suitably prepared watchglass (654).

1311. Examination for K, Li and Na.—Since K and Na have been introduced into the substance by the fusion-mixture, these alkali-metals cannot be tested for in the aqueous solution obtained after fusion.

Unless the analyst is satisfied that the spectroscopic test, made with the special precautions mentioned in paragraph 67, is conclusive, he should proceed to examine the substance for K, Li and Na by paragraph 1313.

The simpler method of examination by paragraph 1315 may be employed if the substance is a silicate: and if K, Li and Na are present in a condition in which they are insoluble in acids, they will usually be contained in a complex silicate.

ANALYSIS OF SILICATES.

1312. The Presence of Silica in a substance will have been shown by the special test (1212) or by VIII. in paragraph 1307.

• When silica has been found it becomes necessary to examine the substance for all metals, since many silicates, which are soluble when they are alone, become insoluble when they are mixed or combined with other insoluble silicates.

Many silicates may be entirely decomposed by heating them with strong HCl for some time just short of boiling. If the decomposition has been complete, only a colourless residue of silicic acid will remain, which may be identified by its insolubility in fused microcosmic salt (648), and by causing effervescence when it is fused into a bead of Na₂CO₃ (649).

If it is found that the silicate is not completely decomposed by hot strong HCl, it should be treated with fused alkaline carbonates, as is directed in paragraphs 1308, et seq. In the examination which follows the fusion, it must be remembered, however, that all metals may be present, and that the metals Na and K must be tested for specially in the original substance by paragraph 1313 or 1315, since they have been introduced in the fusion-mixture.

Examination for K, Li and Na in an Insoluble Substance.

Since Na and K cannot be tested for in the solution which is obtained after fusion with alkaline carbonates (1308), a separate portion of the original substance must be examined for these metals by one of the two following processes. The materials which are used in the tests must be perfectly free from K and Na.

1313. Decomposition by means of CaO.—The finely-powdered substance is mixed with its own weight of sublimed and crystallised NH₄Cl in powder, and with eight times its weight of pure CaCO₃ (1314). The mixture is gently heated in a platinum crucible for a few minutes, and is finally kept at a bright red heat for twenty or thirty minutes. The mass will not fuse as a whole, but sufficient CaO will be dissolved by the fused CaCl₂, and thus be brought into contact with the silicates, to secure their decomposition.

The cool substance is turned out of the crucible if possible, and is boiled with water for some time after it has crumbled by the slaking of the CaO. The liquid is then filtered, and Am₂CO₃-solution is added to the filtrate until it causes no further precipitate of CaCO₃. The filtrate from this precipitate is evaporated considerably, and is then freed from traces of Ca by the addition of Am₂C₂O₄.



1317. Boil a portion of the substance for several minutes with NaOH-solution in a porcelain dish, in order to convert the various cyanides into soluble sodium salts. Since this solution may contain the metals Al and Zn, add Na₂CO₃-solution as long as it causes any precipitate, and boil again for several minutes: then filter, make the cold filtrate just acid with HCl, filter if necessary, and test separate portions as follows:—

Reagent added.	Result.	Inference.
I. FeSO ₄ -solution, freshly prepared 2. FeCl ₃ -solution 3. Add ZnSO ₄ -solution as long as it causes any precipitate The precipitate produced by ZnSO ₄ should be filtered off and fused into a colourless borax bead.	Blue precipitate White precipitate Blue precipitate Blood-red coloration seen on filtration Light brown precipitate White precipitate A blue bead is produced	Presence of ferro- or ferricyanide. Probable presence of cobalticyanide. Presence of (FcCy ₆)iv. Presence of (CNS)'. Presence of (FeCy ₆)'''. Presence of (FeCy ₆)'''. Presence of (CoCy ₆)'''.

- 1318. If a Simple Cyanide only is present, the ordinary course of analysis (1229 et seq.) is pursued, but it will be necessary to remove the HCN by boiling the solution for some time after adding an acid (1229, 1236).
- nary tests (1317), it may usually be decomposed completely, as is proved by testing a portion with FeCl₃-solution, by pouring on the substance which is to be examined for metals some strong INO₅ in a porcelain dish and boiling down nearly to dryness. The liquid is then diluted and boiled, and may be considered as Schriffen III. (1229), any undissolved residue being treated as is directed.

The less simple methods in paragraph 1321 may be employed

1320. If the Presence of Ferro-, Ferri-, or Cobalti-cyanide has been proved by the preliminary tests (1317), two methods of procedure are open to the analyst.

But the cyanogen may be removed from the substance before the analysis (1321); or the substance may be examined such preliminary treatment (1322).

that it enables the analyst to decide whether the metals in complex cyanogen acid-radicles or not. It also precise knowledge of the constitution of the substance.

METHOD I.

may be effected by either of the following processes, 1 or 2. The substance may then be examined for metals in the usual way (1235).

1. Pour upon the powdered substance strong H₂SO₄ in a porcelain crucible, evaporate to dryness and heat the residue until the evolu-

METHOD II.—FOR

1322. Boil the substance with water, filter, and wash the residue

Residue: boil in a porcelain dish with KOH-solution for several minutes. Filtrate: examine this for metals, more especiboil again; filter and wash the residue :ally the alkali-metals, and for acid-radicles, according to the direc-Filtrate: pass H2S, and if it causes any precipitate continue to pass the tions given for analys-ing a liquid (1213). The acid-radicles should filter :be first tested for in a Precipitate: wash well with boiling water, and boil the precipitate with strong HNO₃, a black residue of HgS may remain, filter this off after diluting the acid, and Filtrate: add dilute HNO. portion of the solution, (FeCy₆)iv, (CoCy₆)" and if (FeCy₆)", recipitate: boil with KOH, filter off any black precipitate and examine it for Hg by heating it with Na₂CO₃ one or more are found, Precipitate: confirm the presence of Hg in the precipitate by heating it with Na₂CO₃ in a bulbthe solution must be evaporated to dryness with HNO₃, and the residue strongly heated (General Table 1235) tube (140) or by par. 1250.
The filtrate (or solution if HgS is absent) is evaporated to dryness, the residue is then dissolved in a little hot strong HCl, the in a bulb-tube (140), or by par. 1250.
Add to the filtrate or solution HCl until it is acid, pass H₂S to saturaafter filtering from the solution is much diluted, saturated with H₂S precipitate, in order H2S, and filtered :to destroy these cyano-gen radicles. tion, and examine for As, Sb, Sn, (Pt, Au), by Table II., commenc-The Fe, Co, etc., in the cyanogen acid-radicles will then be detected in Precipitate: examine Filtrate: for Pb and Cu by AmCl, AmOH the General Table (1235). Table II. (1253). in excess, ing at par. 1254. and Am2S, and examine for Zn, Mn. Co, etc. (1296).

Note.—If H₂S causes a precipitate, NaHS or KHS may be added, drop by dro be done instead of passing H₂S to saturation and then adding KOH.

Not Break

tion of H₂SO₄-fumes cease. When the residue is cold, heat it with a little strong HCl, then add water and heat again until all is dissolved, except possibly BaSO₄ and SrSO₄ which are found by par. 1245.

2. Heat the substance strongly in a porcelain crucible after mixing it with three or four times its weight of a mixture of three parts of Am₂SO₄ and one part of AmNO₃.

METALS IN CYANOGEN COMPOUNDS.

with boiling water, adding the washings to the filtrate:-

then add Na ₂ CO ₃ -solution s	s long as it causes an	y precipitate in the s	olution or the filtrate, and				
gas until the liquid is satura	gas until the liquid is saturated, then add more KOH (Note), heat and						
gradually until the liquid is j	ist acid, pass H ₂ S to sa	turation and filter:—	(1235). Fe, Co, Mn, Cr, if], found in this residue, were not present in				
Filtrate: Divide into two par	ets, a, b :		cyanogen acid-radicles.				
a. Examine for acid-radicles in the usual way, testing for (CoCy ₆)" by adding excess of ZnSO ₄ ,	b. Evaporate to dr residue: when col and filter:—	yness and fuse the d, boil it with water					
filtering and trying whether the precipitate gives a blue bead with borax.	Residue: dissolve in HCl, and test for Al, Fe, Mn, Co (1296); the last three if found were present as cyanogen acid-radicles.	Filtrate: acidify a portion if yellow, with HA and add PbÅ ₂ , a yellow precipitate shows (CrO ₄)", the Cr having been present as cyanogen acid-radice. Test another part	. ,				
	,	for Al by adding HCl in excess, then AmOH in slight excess.					

until it no longer gives any precipitate in the liquid or in the filtrate; this may

This table rnns ACTORS both pages.

1323. GENERAL GROUP TABLE FOR

Group I .- Reagent HCl.

Group II.-Reagent H2S.

The Group-Pp. may contain in addition to AgCl, Hg₂Cl₂, PbCl₂:—

TICI (white). H_2WO_4 (,,).

TICI will usually be readily detected by 201 will usually be readily detected by a spectroscopic examination of the original substance or of the group-pp. It may be removed from the group-pp. by boiling with a little water, and ppd from the cold filtrate by KI (Pb is also thus ppd'): the pp. is yellow, and gives the thallium spectrum.

H₈WO₄: a white pp. becoming yellow when the liquid is boiled. Its presence is confirmed by dropping a piece of Zn into a portion of the acid liquid and pp., when a deep blue colour will be obtained.

Also by fusion of the pp. in a microcosmic bead, which will be colourless or yellow in the outer flame, blue (or with FeSO₄, blood-red) in the inner flame.

The Group-Pp. may contain as sulphides, in addition to sulphides of Hg, Pb, Bi, Cu, Cd:—

Pd-sulphide (black-brown)
Tl-sulphide (black)
[Os, Rh, Ru*-sulphides] Insoluble in

And in addition to SnS, SnS₂, Sb₂S₃, As_2S_3 , Au_2S_3 , $PtS_2 : -$

Am₂S.

[Ir-sulphide*] (brown) Soluble in Se-(red-yellow) (black) Am2S. Te-,,

The yellow colour of a solution containing V is changed to blue by H.S.

TI is readily found by the spectroscope in the original 24. is readily found by the spectroscope in the original substance or in the group-pp.

Pd will remain in solution in excess of AmOH with 2, Cd and Cu in Table II.a. (1253): from this solution it is ppd by adding HCl: its presence may be further confirmed by the HgCy₂-test (213), or the KI-test (214).

The examination of the sulphides in Group II.B., when the above elements may be present, is best commenced by fusion with Na₂CO₃ and Na₂O₂ or NaNO₃. From the cooled mass, water dissolves sodium-arsenate, -molybdate, -selenate, and -tellurate, leaving SnO₃, sodium-antimonate, Au, Pt, and Ir undissolved; the elements may then be detected by special tests. For the separation and detection of the platinum metals a special work must be consulted a special work must be consulted.

* Only completely ppd. If the liquid has been warmed, and H_2S passed for a long time.

PRECIPITATION OF RARER ELEMENTS.

Group III.—Reagents AmCl, AmOH, and Am₂S. Precipitated from the filtrate from Group III. by excess of HCl. Group V. The Group-Pp. may contain as sulphides, in addition to Fe, The filtrate, after the separation of all the group-precipitates, may contain besides Mg, K, As sulphides :-V (black). Zn, Mn, Ni, Co:-W (trace, brown).
Ni (trace, black). U (black brown.)
In (, , ,)
Tl (,, , ,) and Na:— Li, Cs, Rb. And as hydroxides in addition to Al, Cr (and phosphates, etc.):— This pp. is dried and then fused with a mixture of Na₂CO₃ and KNO₃: on boiling the mass with water, NIO remains undiscalued. These metals are readily detected by their very characteristic spectra (66).

Before proceeding to the spectroscopic examination, it is well to remove the Mg and to convert the alkali-metals into chlorides by the procedure given in the second paragraph of Note 3 (7253).

Pour a few drops of HCl upon the residue thus obtained, and evaporate the solution almost to dryness; triturate the residue with 90 per cent. alcohol, and filter. The alcoholic filtrate will contain all These metals are readily de-Ti (white, flocculent). Be (colourless, gelatinous). NiO remains undissolved; it is filtered off, and its presence confirmed by fusion [Zr [Ta, Nb, Ce, La, Di, Yt, E, Th.] in a borax bead (313). he aqueous solution may contain alkaline vanadate and tungstate. V is sepa-rated by saturating the liquid with AmCl, and its presence confirmed in the pp. Tl and In will usually be readily detected by a spectroscopic examination of the pp. examination of the pp.

77 may be first separated and
detected by dissolving some
of the group-pp. in boiling
dilute HCl, and reducing any
Fe present in the solution by
H_SSO₃; then precipitating Tl
by addition of KI, and testing
the pp. by the spectroscope. presence confirmed in the pp. by the borax bead (397), and by dissolving the pp. in HCl and employing the reactions with H₂O₂ and with Zn (396, 393).

(The filtrate, after concentration if necessary, is acidified with HCl, and the presence of W confirmed by the Znthe Li, Rb and Cs, with small quantities of K and Na. If traces of Rb and Cs only are present, evaporate and extract with alcohol several times. Evaporate the alcoholic extract A more complete scheme for the analysis of this group-pp. will be found in par, 1324. of W confirmed by the Zn-reaction and the micro-cosmic bead (128, 131) to dryness, add a little strong to dryness, add a little strong HCl, evaporate once more, gently ignife, and triturate the residue with equal parts of anhydrous ether and absolute alcohol: then filter through a dry filter. The filtrate which contains the Li may be evaporated to dryness and tested by the fiame coloration and the spectroscope (66). by the fiame coloration and the spectroscope (66). The residue is dissolved in a little HCl, and K, Rb and Cs are ppd by H_pPtCl₀: this pp. is boiled with small quan-tities of water until it no longer gives the K spectrum, the spectra of Rb and Cs will, if present, he given by the if present, be given by the

TABLE FOR ANALYSIS OF GROUP III. PRECIPITATE, 1324. The precipitate may contain Fe, Tl, Al, Cr, Zn, Mn, Ni,

These lines run cross both ages.

After a preliminary spectroscopic examination has been made of a small portion of the pp. for Tl and the group pp. is dried and fused in platinum with KHSO4 for some time, the cold mass is then pow-

Residue: may contain Ta₂O₅, Nb₂O₅, also SiO₂, and traces of Fe and Cr which have escaped solution.

By fusion with KClO₃ and NaOH, Cr and Si are rendered soluble in dilute NaOH; from the residue, insoluble in NaOH, Ta₂O₅ and Nb₂O, may be removed by washing with dilute Na₂CO₃solution.

Solution: reduce Fe by adding $\rm H_2SO_3$ in excess, dilute considerably and passing a stream of $\rm CO_2$ unless it smells constantly of $\rm SO_2$:—

Pp. may consist of TiO_2 , possibly also traces of Zr.

Confirm the pre-sence of TiO2 by the microcosmic bead (339).

Filtrate: add a few drops of strong HNO3, concen-more by adding excess of AmOH:—

Pp. may contain Fe, In, U, Cr, Al, Be, etc. Dissolve

Pp. may contain Fe, In, U, etc. Dissolve in HCl, of HCl, add BaCO3, and let stand :-

Pp. may contain Fe, In, U, also traces of Al and Cr. It is dissolved in HCl, and NaHCO₃ is added in excess, whereby U is obtained in solution. In is found by the spec-

troscope.

Filtrate may contain Zr, Ba is ppd by adding liquid is concentrated, with KOH. Crystals the liquid is allowed The pp. is washed with

Pp. may contain Zr,
Th, Ce. Th and Ce are dissolved away by boiling with dilute HCl, and are reppd by AmOH.

WHEN THE RARER ELEMENTS MAY BE PRESENT.

Co, Ta, Nb, E, Be, Ti, Ce, La, Zr, Th, Yt, In, U, Di.

In, and Tl has been further tested for, if necessary, by KI as directed in the General Table (1323), dered and allowed to stand, with shaking, for a long time with cold water:—						
boil for a long time, preser	ving the liquid from oxidation by the air by covering the	ne vessel, and by				
trate by evaporation, and a	dd AmOH in excess: filter, dissolve the pp. in HCl, and	I precipitate once				
it in HCl, and add excess of	it in HCl, and add excess of cold strong KOH-solution :— Filtrate may contain :—					
boil off any large excess	Filtrate may contain Al, Cr, Be: dilute and boil for some time, filter. Al remains in solution, and may be ppd by adding HCl just in excess, then AmOH	Zn, Mn, Ni, Co.				
Ce, La, Di, Yt, E, Th. excess of H ₂ SO ₄ , the and exactly neutralised of K ₂ SO ₄ are added, and to stand for some hours. K ₂ SO ₄ -solution.	in excess. Cr and Be are separated by fusion with Na ₂ CO ₃ and KClO ₃ , dissolving in water and pps, Be by adding HNO ₃ in excess, then AmOH in excess.					
Soln. may contain Y, E, also Be. The solution is ppd by AmOH, and Be dissolved away from it by H ₂ C ₈ O ₄ .						

GENERAL METHODS FOR DETECTING ORGANIC SUBSTANCES.

INTRODUCTORY REMARKS.

1330. The Detection and Identification of an Organic Substance is usually more difficult than the detection of an inorganic substance, and the difficulty is increased when a mixture of organic substances has to be dealt with.

Since no complete scheme of separation is possible, recourse must be had to several partial separations, such as the precipitation of one or more of the substances in an insoluble form, or the use of a suitable liquid which is a solvent for certain substances but not for others. Sometimes separation may be effected by the application of heat, the more volatile substance passing off as vapour and leaving the non-volatile substance behind. Fractional distillation (672) may be utilised for the separation of liquids of different boiling-points; distillation in steam and in vacuo may also be resorted to.

The suggestions and Tables which are given below include:-

- (1) Preliminary tests, such as the action of heat, acids and other reagents.
 - (2) The identification of the elements present.
- (3) The identification of the compounds by their special properties and tests.

The student should be familiar with the methods of detecting the elements in an organic substance (673-688), and with the reactions and properties of organic substances (689-1148) before he applies them in the following scheme.

SCHEME FOR THE DETECTION OF ORGANIC SUBSTANCES.

PRELIMINARY EXAMINATION.

1331. The Physical Properties of the Substance, such as its colour and smell, are noted; also its physical condition, and, if solid, whether it is amorphous or crystalline.

The determination of its solubility in various solvents will frequently indicate the chemical character of the compound, since:—

Organic salts are usually soluble in water, but insoluble in ether. Organic acids are often soluble in hot water, but sparingly soluble in alcohol and in ether.

Hydrocarbons and their haloid derivatives are insoluble in water, but soluble in ether.

Alkaloids and organic bases are usually insoluble in water, but soluble in acids and in organic liquids.

The Table of Solubilities (1336) will be of use after trying the action of solvents.

A knowledge of the melting-point (669), boiling-point (671), and density (668) is also often of great assistance.

Note also the effect produced by the substance or its solution on litmus. Acid reaction usually indicates the presence of free acids, or of hydrolysed salts of certain bases; esters also usually contain free acid. Many alkaloids and basic substances have an alkaline reaction to test-paper; but a large number of organic bodies such as the hydrocarbons, alcohols and ethers are neutral in reaction.

1332. Ascertain what Changes the Substance undergoes when it is heated in a Test-tube, whether it is volatile or fusible, and whether any gases or vapours are evolved.

EFFECT OF HEATING THE SUBSTANCE IN A TEST-TUBE.

	IDATING THE SUBSTANCE IN A	
Result.	Observation.	Inference.
a. The substance volatilises, leaving no residue.	The vapour is non-inflammable. The vapour burns with a pale blue flame. The vapour burns with a blue flame. The vapour burns with a pale blue flame; slight charring occurs. The vapour burns with a smoky flame. The vapour burns with a luminous flame. The vapour is acid, pungent, and inflammable.	Chloroform. Alcohols, aldehydes. Chloral, carbon disulphide. Glycerol. Aniline, benzene, naphthalene, turpentine. Ether, ethyl acetae, benzaldehyde, ni- trobenzene. Formie acid, acetic acid.
b. The substance fuses, and volatilises without residue.	The vapour burns with a smoky flame. NH, and CO ₂ formed. Pungent fumes yielding a white sublimate. Pungent fumes, burning with a smoky flame; and a sublimate is formed. A slight explosion.	Phenol. Urea. Oxalic acid, succinic acid. Benzolc acid. Pieric acid.
c. The substance fuses and chars.	It chars slowly, evolving CO. It chars slowly, evolving acetone. It emits a smell of charred sugar.	Formate. Acetate. Tartrate, citrate, malate, meconate, pyrogallol, sugars.

Result.	Observation.	Inference.
e. The substance neither chars nor fuses.	A carbonate remains (474).	Oxalate.
f. The substance fuses to a coloured liquid.	The liquid is reddish; vapour of un- pleasant smell is emitted, which burns with a smoky flame.	Hippurate, or an alkaloid.
g. The substance fuses, and evolves cyanogen.	No further special observations, A smell of H ₂ S is also noticed. The substance darkens in colour.	Cyanide. Thiocyanate. Ferrocyanide, or ferri- cyanide.

1333. Heat the Substance with free exposure to the Air on a piece of thin porcelain, or on a nickel spatula: if a black residue of carbon is left, burn off the carbon by heating it strongly from beneath in the blowpipe-flame (Exp. 21, page 39).

One of the following results may be obtained:-

- (a) A Coloured Residue may remain and will probably be due to the presence of a metal which produces a coloured oxide.
- (b) A White Residue may remain, which when it is moistened with water shows an alkaline reaction to test-paper: this may be caused by the oxide or carbonate of K, Na, Ba, Sr, Ca, or Mg derived from an organic salt.

The residue, in either case, may be examined by the preliminary tests 1219, 1220, and may then be dissolved (1229) and examined by the General Table (1235).

- 1334. Mix the Substance with powdered Soda-lime and heat, noting carefully the characteristic smell of any of the following substances, which may be also recognised by special tests:—
 - (a) Ammonia (426), is evolved from nitrogenous substances, such as ammonium-salts, urea, alkaloids, urate and hippurate.

Nitrocompounds, such as nitrobenzene and picrate, do not give off ammonia when they are heated with soda-lime.

- (b) Aniline (978), evolved from aniline compounds.
- (c) Benzene (814), evolved from benzoate or hippurate.
 - (d) Phenol (848), evolved from salicylate.
- 1335. Try the Action of Strong Sulphuric Acid upon the Substance, both in the cold and on heating: the results which may be obtained are tabulated below:—
- Mis test may be supplemented by heating the substance with dilute H₂SO₂, since formates are tates and cyanogen-compounds may be frequently detected by this means.

Effect of adding Strong $\mathrm{H}_2\mathrm{SO}_4$ to the Substance, or to its Strong Solution, and then Heating.

Result.	Observation.	Inference.
a. No change occurs in the cold, and either little or no charring on heat- ing.	CO evolved, no charring. HA evolved, no charring. CO and CO ₂ evolved, no charring. Pungent fumes, no charring. Slowly darkens; irritating fumes evolved. Slowly darkens; acid fumes, then CO and CO ₂ evolved. (Difference from tartrate, etc.) Slowly darkens; CO and CO ₂ evolved. (Difference from tartrate, etc.) Slowly darkens; bungent fumes evolved. Effervesces, slowly darkens. HCN evolved, no charring. HCN and finally CO evolved, white precipitate. CO evolved, and liquid becomes turbid. SO ₂ evolved, yellow S formed. Slight charring.	Formate. Acetate. Oxalate. Chloral; chloroform. Succinate, or benzoate. Malate. Citrate. Urea. Urate. Cyanide. Ferricyanide. Ferricyanide. Thiocyanate. Meconic acid.
b. No change occurs in the cold, but the substance turns dark-coloured, or chars, when it is heated.	Fragrant smell; vapour burns with a smoky flame; SO ₂ is evolved, and charring occurs. Acrolein is smelt, and ultimately SO ₂ is evolved with much charring. CO and SO ₂ are evolved, with gradual charring. CO, CO ₂ , and SO ₂ are evolved, with rapid charring. SO ₂ is evolved, and charring occurs. The substance then turns reddish-brown, evolving CO ₂ and pungent fumes. The substance turns brown, and SO ₂ is evolved. A vivid red coloration is produced, and SO ₂ is evolved. Characteristic smell, then charring, and evolution of SO ₂ .	Alcohols. Glycerol. Most sugars. Tartrate. Hippurate. Salicylate. Tannate. Gallate. Turpentine.
c. The substance usually becomes coloured in the cold; the colour deepens on heating.	Colourless or pale brown, cold; blackening on heating. Yellow, cold; puce, on heating. Pink, cold; blackening on heating. Colourless, cold; blue on heating. Colourless, cold; blue on heating. Pink, cold; violet, then brown on heating. Pink, cold; violet, then brown on heating. Yellow, cold; brown-red, then violet on heating. Yellow, cold; red on heating. Blood-red coloration, cold. Yellow, brown and finally red coloration. Forms white sulphate, soluble in excess. Yellow, cold; charring on heating, and evolving CO and SO ₂ . Brown, cold; charring rapidly, and evolving CO and SO ₂ . Solid cane-sugar gives these changes slowly.	Quinine, cinchonine, strychnine, etc. Narcotine. Brucine. Codeine. Papaverine. Morphine. Narcotine. Veratrine. Salicin. Digitalin. Aniline, pyridine, quinoline. Starch. Cane-sugar.

1336. Test if the Substance is Soluble in or Miscible with :-

(1) Cold water, (2) Hot water, (3) Alcohol, (4) Ether.

The knowledge of the solubility of the substance may aid in its detection, and may further enable a separation of mixed substances to be effected by the use of suitably selected solvents.

TABLE OF SOLUBILITY OF ORGANIC SUBSTANCES.

s = soluble. i = insoluble. ss = slightly soluble.

vs = very soluble. vss = very slightly soluble.

						Wat	er.		
	Substan	e.				Cold.	Hot.	Alcohol.	Ether.
Acids:—									
Aliphatic:				_		vs	vs	vs	i
Formic .		•	•			vs	vs	vs	S
xalic .						s	VS	VS	s ·
Succinic						SS	VS	s vs	VSS
Malic .						vs	* VS	S	i
Tartaric				•		s s	S	6	î
Citric .				•	.	ь	۵		-
Aromatic:						ss	s	s	s
Benzoic.		•	•			88	S	s	vs
Salicylic		•	•	•	: 1	SS	s	s	8*
Gallic .		•		•		S	s	SS	VSS
Tannic -		•	•	•	: 1		8	8	SS
Hippuric		•	•	•		8S	s	S	vss
Meconic		•	•		.	VSS	VSS	SS	i
Uric . Hydrocarbons	AND DE	RIVAT	IVES:						
Turpentine	5, AND 1016	241 1 12-			. }	i	i	vs	VS
Benzene							i	8	s
Naphthalene	•					i	i	vs (hot)	s
L'apronaiche			-			_	1 .	\ vss (cold)	
Anthracene						i	i	88	SS S
Chloroform	<i>7</i> .					i	i.	s	ь
Carbon tetra		· .				· i	i	71	s
//odoform ~	r					î	i	vs	vs
Carbon disul	phide Y					i	i	. VS	1
ALCOHOLS, ET	HERS:—					vs	vs	vs	vs
Methyl alcol				•		vs	vs	vs	vs
Ethyl alcoho)i		•	•	•	vs	VS	vs	vs
Amyl alcoho		•	•	•		8	8	s	i
Glycerol (Gl		•	- *	•	•	88	88	vs	vs
Ethyl ether		•	•	•	•	VS.	vs	VS	VS
Ethyl aceta	ie	•	· •	•	•	7			
Phenoi (Car	halfa acid	1				8	8	vs (hot)	8
Hydroguino		,				S	g.	s	į s
Pyrogaliol (Pyrroga Ilio	hise	٠.			vs	VS	vs	s
Trinitrophe	rol (Pieri	acid	3			VSS ·	s	S	S
AEDEHYDES,			y -				j et		
Formaldeh				· .	-	SS	. 88	8	i
Acetaldehv	le.			7 .		vs	VS	S	8
Trichloraid	hyde (Ch	loral)				S	8	s	
Benzaldehy	de .			1.5		S	8	VS	vs
Acetone (D	imethyl k	etone		. `-		8	· 8	· 8	s
GAR BOHYDRA	PES:-	*				1		8.	s(CH s.OH
Dextrose (rape sug	ar) .	* j		•	VS	VS	* 8	3(0118.011
Laevulose (Fructose)		· ; , *	Y - 1 0	- T =	\$ 576	- 8	SS	i
C Sucrose (Ca	ne sugar)	1 . 1		شي آج		vs s	VS.	8	i i
Malfose (M Eactore (M ICallulose Stanch Dexistra	ait sugar)		g 🔑 🍃	1750	44.	eé .	88	vss	î
Lactore (M	uk sugar)	300	ā-y °, ,•		Sec.	380	i	i i	Î
Centrose	114	300	g in the	145 .	1200	1 1 10 1	8	i.	1. 1
STREET		157	- 37, 3*	1.50			, , , , , , , , , , , , , , , , , , ,	i	, i
Dexum.					3 1 3		e de la		1
· · · · · · · · · · · · · · · · · · ·	五五二世三月	2.00		H . W -	1 1 1 1 1	. M. 1894 . 3	. 1		

	~ .						Wa	ter.		
	Sui	ostano	e.				Cold.	Hot.	Alcohol.	Ether.
GLUCOSIDES :- Salicin . Digitalin							VSS	s	s (hot)	·
Amygdalin Organic Basi	: ES:—	:	:	:	:	:	SS S	ss	s (hot)	vss i
Aniline . Toluidine	:	:	:	:			VSS SS	VSS	s vs	S
Pyridine Ozinoline					÷	:	miscible	miscible	_	_
Acetanilide	Antif	ebrin)	:	:	:	:	VSS SS	VSS SS	s i	8 1
Phenacetin Antipyrin (I	henaz	(are)	٠	•	•		i s	VSS S	ss	i
Urea (Carba	mide)		·	:	:	:	vs	vs	s vs	SS SS
5 Quinine.							i	i	vss	SS
Ginchonine Morphine	:	:	:	:		•	i vss	i Vss	SS S	VSS VSS
2 Codeine. Narcotine	•						s	8	vs	i
Papaverine	:	:	:	:	:		i	i	SS SS	s i
Strychnine Brucine.	:	:	:	:			i ss	i ss	s (hot)	i
3 Caffeine (Th	eine)				:		SS	S	SS	vss
Veratrine	:	:	:		:	:	ss i	ss i	s s	8 8
 Atropine Hyoscyamin 	e.	:	:	:	:	:	iss	i ss	vs vs	8
Hyoscine Nicotine	•	•	•	•			88 *	SS	S	8
3 Coniine		:	:	:	:	:	VS SS	vs s	vs vs	ys vs

SPECIAL EXAMINATION OF ORGANIC SUBSTANCES.

1337. If the substance is a simple one, it will probably have been identified by the preceding preliminary examination, and it will then be at once submitted to confirmatory tests.

But if the preceding treatment has given any indication that the substance is a mixture, some method of separation should be devised. The action of solvents (1336), or treatment with solutions of acids or of alkalis may be tried upon a solid substance. A liquid may be subjected to fractional distillation (672).

In any case it is advisable to ascertain what elements are present in the substance: the presence of the more important elements will be indicated by the following experiments.

- 1338. Heat the dry Substance strongly in a Test-tube with dry CuO (674, 676), and observe the results:—
- (a) CO_2 is evolved (474), whereas the original substance evolved no CO_2 when it was treated with dilute HCl, indicating the *Presence* of C.
- (b) Drops of water condense on the inside of the tube, indicating the Presence of H.
- 1339. Heat the Substance with a little Sodium in an ignitiontube until it is charred, and drop the tube into a few c.c. of water contained in a mortar: then grind up the fragments with the

water, filter, and divide the filtrate into three portions:-

(a) To the first portion add a small crystal of $FeSO_4$, boil the liquid and acidify it with HCl: a blue or green coloration shows the *Presence of N* (679).

- (b) In the second portion dissolve a small crystal of sodium nitroprusside: a violet or purple colour shows the *Presence of S* (681).
- (c) If the substance contains no nitrogen, add to the third portion HNO₃ and AgNO₃-solution: a white or yellow precipitate shows the *Presence of Cl., Br., I* (688).
- (d) If the substance contains nitrogen, ignite a fresh portion with soda-lime free from chloride in a combustion-tube. Shake out the contents of the tube into dilute nitric acid, filter the liquid and add AgNO₃-solution; a white or yellow precipitate shows the Presence of Cl, Br, I (688).
- 1340. Mix the charred Substance with Mg-powder, heat the mixture strongly and moisten the cold residue: an onion-like smell indicates the *Presence of P* (684).
- 1341. Hydrolysis may be resorted to for the identification of certain substances, such as esters, alkyl cyanides, glucosides and sugars, which may by this means be split up into readily recognisable products. The process consists in the prolonged heating of the substance with a moderately strong acid, such as HCl or H₂SO₄, or with KOH-solution. The process may be carried out in a flask connected with a reflux condenser.

The resultant products include alcohols, glucoses and organic acids, and these may be detected by the methods given under the separate tests and reactions for these substances.

1342. The Presence of an Alkaloid may be ascertained by the application of the general tests given in paragraphs 1036–1042. The identification of the particular alkaloid present may then usually be effected by carefully extracting with warm pure alcohol acidified with dilute HCl, driving off the excess of alcohol, diluting the extract with water and the graphs 1050–1148.

A more general method for separating the alkaloid consists in making the mixture alkaline with NaOH-solution, evaporating the whole to dryness, and extracting the residue several times with small quantities of ether. The ether is then distilled off, and the residue consists of the alkaloid free from inorganic and from most organic substances (Note). If aniline, pyridine or quinoline are present it will be necessary to remove them by steam distillation:

Note: Morphine is insoluble in ether; but this alkaloid may be extracted to the fact anyl alcohol from the residue remaining after the extraction by ether

1343. The Presence of Organic Acids or their Salts in solution can usually be determined by the formation of characteristic insoluble compounds on the addition of solution of a salt of iron, calcium, or silver (1344, 1345, 1346).

The organic acid-radicles should be present in sodium, potassium, or ammonium salts, since other metals interfere more or less seriously with the detection of the acid-radicles by these tests. The solution is accordingly prepared as is directed below, under a, b, c :

(a) If alkali-metals only are present, the solution to be tested must be rendered neutral or <u>very faintly alkaline</u>. The most suitable reagents for producing this condition are dilute HNO₃ and NaOH-solution. According to the reaction of the solution, one or other of these reagents is added until the solution does not affect the colour of test-papers.

(b) If metals other than the alkali-metals are present, the substance must be boiled with excess of Na₂CO₃-solution, filtered, and acidified with HNO₃; the CO₂ is then boiled off and the solution is constally particularly with Na OH.

tion is carefully neutralised with NaOH-solution.

(c) If metals of Group II. or III. are present, the following treat-

ment may be requisite to ensure their removal:-

The solution is acidified, if it is not already acid, with dilute HCl, and H₂S is passed to saturation; the liquid is then filtered, and AmOH and Am₂S are added to the filtrate: the liquid is then filtered again and the filtrate is reserved. The filtrate will contain the ammonium salts of the organic acid-radicles, together with excess of Am₂S, and its further treatment will depend upon whether only Ba, Sr, Ca, Mg are present, or other metals are also present:—

If no other metals are present, acidify the liquid with dilute HCl, boil off the liberated H₂S, filter if necessary, and neutralise the filtrate with NaOH-solution. This liquid may now be tested with the reagents mentioned in pars. 1344, 1345. (See Note below.)

If other metals are present eidify with dilute HCl, boil off the H₂S, add Na₂CO₃-solution excess, filter, acidify with HNO₃, boil off CO₂, and carefully neutralise with NaOH-solution: then proceed to test the liquid by pars. 1344, 1345.

Note.—A separate solution of the original substance must be prepared for the AgNO₃ test (1346), in which dilute HNO₃ is used instead of HCl, else the AgNO₃-solution will give a white precipitate of AgCl which may be mistaken for an organic salt.

The results given by the tests in pars. 1344-1346 are not pronounced, and are in some cases not even visible, if the reagents are added to very dilute solutions, and strong solutions should therefore always be employed.

It will be seen that the results which are yielded by other substances besides the organic acid-radicles are given in paragraph 1344.

1344. Action of FeCl₃-Solution on Neutral Solution of Organic Substances.

Result.	Observation.	Inference.
a. A red coloration is produced in the cold.	The coloration is destroyed by HgClls- solution, and is unaltered by dilute HCl. The coloration is destroyed by dilute HCl; a red precipitate forms on boiling. (Diff.) In the original solution AgNO3 gives a black precipitate in hot solution. (Diff.) Strong H2SO4 gives a smell of HÅ. The coloration is with difficulty destroyed by HCl. CaCl2 produces a white precipitate in the original solution on boiling. (Diff. from formate and acetate.) The coloration is destroyed by HA, but not by HCl.	Thiocyanate. Formate, or acctate. Formate. Acctate. Meconate.
b. A purple colora- tion is produced in the cold.	The purple colour is not destroyed by HĀ. A salicylate when heated with strong H ₂ SO ₄ and CH ₃ OH gives off methyl salicylate. (Diff. from phenol.) The purple colour is destroyed by HĀ. Phenol solution when treated with AmOH and bleaching-powder turns blue. (Diff. from a salicylate.)	Salicylate. Phenol.
c. A blue - black precipitate is pro- duced in the cold.	The precipitate disappears on boiling. With a gallate KCN produces a red coloration. (Diff. from a tannate.) The precipitate is permanent on heating. Ammoniacal CuSO ₄ -solution produces a green precipitate with a tannate. (Diff. from a gallate.)	Gallate.
d. A blue precipitate is produced in the cold.	The colour of the precipitate is changed by KOH into brown.	Ferrocyanide.
e. A buff or reddish precipitate is pro- duced in the cold.	The addition of HCl gives crystalline scales of benzoic acid. The original substance burns with a smoky flame. (Diff. from a succinate.) The precipitate is soluble in HCl. A succinate is precipitated on the addition of BaCl ₂ , AmOH, and alcohol. (Diff. from benzoate.) On the addition of HCl. appurio acid	Benzoate. Succinate.
	On the addition of HCL suppurio acid separates. The original substance burns with a smoky flame, and evolves NH, and C ₂ H ₂ when it is heated with sode-lime. The reddish precipitate dissolves in HCl. Confirm by adding to the original substance K ₂ Cr ₂ O ₇ and H ₂ SO ₄ , a blue coloration appears.	Hippurate. Aniline.
f. A coloration is produced in the cold.	Brown or olive-green coloration. Confirm by adding FeSO ₄ -solution to the original substance, a blue precipitate forms. The coloration is blue.	Ferricyanide. Morphine.
g. No effect is pro- duced until the	The solution reddens on being heated and forms a red precipitate.	Many alkaloids and organic bases.

1345. Action of $CaCl_2$ -Solution on Neutral Solution of Organic Salts.

Result.	Observation.	Inference.
a. A white precipitate forms in the cold.	The precipitate is insoluble in HÅ and in KOH; when dried and heated, it scarcely chars and leaves a carbonate. The precipitate is soluble in HÅ and in KOH; it often only appears on shaking, and chars when dried and heated. Confirm by heating the original solution with ammoniacal AgNOs (759, Note). Add PbÅ2 to the original solution, a white precipitate forms. The precipitate is decomposed by HCl, with separation of uric acid. The precipitate is unchanged when heated.	Oxalate. Tartrate. Meconate. Urate. Hippurate, benzoate, or tannate.
b. A white precipitate forms on heating.	The precipitate is soluble in HÅ and insoluble in cold KOH; it chars on being dried and heated. The precipitate forms more readily in the presence of alcohol; it chars on being heated. Confirm by adding PbÅ ₂ to the neutral solution, and fusing the precipitate under water (758).	Citrate. Malate.
c. A white precipitate forms in the presence of alcohol.	Confirm by adding to the neutral solution FeCl ₃ , a reddish precipitate forms.	Succinate.

 $^{\circ}$ 1346. Action of ${\rm AgNO_{3}\text{-}Solution}$ on Neutral Solution of Organic Salts.

Result.	Observation.	Inference.
a. A white precipitate forms in the cold.	The precipitate is unaltered when heated. The precipitate dissolves when heated: it is not formed in dilute solution. The precipitate forms in strong solutions, rapidly turning black; in dilute solutions a black precipitate of Ag is formed when the liquid is heated. The precipitate turns grey when it is heated. The precipitate slightly darkens when it is heated. The precipitate rapidly turns black. The precipitate is soluble in AmOH, and insoluble in dilute HNO ₃ . The precipitate is is soluble in AmOH, and insoluble in dilute HNO ₃ . White curdy precipitate. soluble in NH ₄ OH and in HNO ₃ . Immediate black precipitate.	Oxalate, meconate, salicylate, malate, or hippurate. Acetate. Formate. Citrate, or malate. Succinate, or benzoate. Tannate, gallate, urate, phenol, and hydroquinone. Tartrate. Cyanide or thiocyanate. Ferrocyanide. Cyanate. Pyrogallol.
b. An orange precipitate is formed.	The precipitate is soluble in AmOH, and • insoluble in dilute HNO ₃ .	Ferricyanide.

EXAMPLE SHOWING HOW TO ENTER THE RESULTS OF ANALYSIS OF A COMPLEX SUBSTANCE.

1347. The substance given for analysis consisted of a powder containing pink, blue, white, and black particles. It smelt faintly of ammonia.

PRELIMINARY EXAMINATION FOR METALS.

Experiment.	Observation.	Inference.
EXPT. I.—Heated in a small dry test-tube.	Water given off which turned turmeric-paper brown. Strong smell of NH ₃ . Substance blackened, no smell of burning. Slight white sublimate.	Pres. of H ₂ O. Pres. of NH ₄ . Pres. of NH ₄ . Prob. pres. of Co, Cu, and abs. of T and A. Pres. of As, NH ₄ , Hg.
Confy.—Held in the upper part of the tube a glass rod with a drop of lime-water hanging on its end. Confy.—Boiled a portion of the substance with KOH-solution. Confy.—Heated strongly another portion of the dried substance with Na ₂ CO ₃ in a bulb-tube.	The lime-water became milky. Brown nitrous fumes evolved, recognised by their smell. Cl evolved, found by its smell and by bleaching litmus. NH, was evolved, recognised by its smell, and by giving white fumes with strong HCl. No mirror formed.	CO ₂ evolved. Pres. of nitrate. Pres. of Cl. Pres. of NH ₄ . Abs. of Hg and As.
Exp. II.—Heated a portion of the substance on a loop of platinum-wire in the Bunsen-flame, moistened with HCl and heated again in the flame. Heated for some time in the fip of the inner blowpipe-flame, moistened with HCl, and again held in the Bunsen-flame.	Bright yellow flame. The flame appeared crimson through the indigo-prism. Crimson coloration, appearing intense red through the indigo-prism. Bright green flame with blue core.	Pres. of Na. Pres. of K. Pres. of Sr. Pres. of Cu.

PRELIMINARY EXAMINATION FOR METALS. (Continued).

Experiment.	Observation.	Inference.
EXPT. III.—Heated a portion of the substance on charcoal in the inner blowpipe-flame. Confy.—Fused in a clear borax-bead in the outer and inner blowpipe-flames. Confy.—Fused on platinumfoil with Na ₂ CO ₃ and KNO ₃ . Confy.—Fused on charcoal in the inner blowpipe-flame with Na ₂ CO ₃ .	The greater part of the substance fused readily, and was absorbed by the charcoal. Deflagration occurred. A red metallic residue remained. In the outer flame a bead green whilst hot, blue when cold. In the inner flame red and nearly opaque. No bluish green or yellow mass on cooling. Red metallic residue. A portion placed on a silver; coin and moistened gave no black stain.	Pres. of a salt of K, Na. Pres of chlorate, nitrate. Pres. of Cu. Pres. of Cu. Abs. of Mn and Cr. Pres. of Cu. Abs. of S.

PRELIMINARY EXAMINATION FOR ACID-RADICLES.

Experiment.	'Observation.	Inference.
EXPT. I.—Added dilute HCl without heating. Heated to boiling.	A colourless gas was evolved, which was free from smell, and turned a drop of limewater milky. Cl was evolved, recognised by its smell and by bleaching moist litmus-paper.	Pres. of carbonate. Abs. of sulphite, hypochlorite, etc. Pres. of nitrate, chlorate, or some other oxidising substance.
Expr. II.—Added strong H ₂ SO ₄ . Confy.—Dropped in copper turnings and heated. Heated strongly, cooled, and rinsed out tube. Confy.—Boiled a portion of the substance with water, added strong H ₂ SO ₄ , cooled and poured in FeSO ₄ -solution carefully.	A bright yellow chlorous gas evolved, which crackled when warmed. Reddish brown fumes evolved. The tube when dry was seen not to be etched. A brown ring formed on the surface of the acid.	Pres. of chlorate. Pres. of nitrate. Abs. of fluoride. Pres. of nitrate.

ſ1347,

These lines run across both pages.

Examination for Metals

Boiled a portion of the substance with FeCl₃, FeSO₄ and KOH,

Boiled a portion of the substance with water; as it did not completely undissolved residue: effervescence occurred, and on boiling, Cl was perceived; the substance was completely dissolved. Cooled this solnto give no pp. on addition of a little dilute HNO_3 :—

No pp.	Diluted with water and p	passed H ₂ S until the liquid smelt strongly of
Abs. of Group I. Hg', Ag, and prob. Pb.	A black pp. Exam ^d . by Table II.	Filtrate which gave no further pp. with H ₂ S Boiled until it no longer smelt of H ₂ S, tion of HNO ₃ (Pres. of Fe). Warmed the some AmHMoO ₄ -solution, warmed; no
	· •	A brown pp. Examd, by Table III.A.
		;.
		25

1347]

IN THE WET WAY.

added HCl in excess, no blue pp.:-Absence of CN.

dissolve, the sol^{n.} was decanted, and a little dilute HCl poured upon the smelt. Added a little strong HCl, boiled as long as any smell of Cl was and mixed it with the water sol^{n.} which had been proved previously

esidue with HCl, it dissolved up.: Absence of (PO_4) . To the	completely (Abs. of Silicate). Ad-	ne soln. changed to light brown on addided a small portion of the HCl soln. treess of AmOH, and boiled; filtered:
A black pp. Examd, by Table III.B.	Filtrate was yellow (Absence	of Ni): added Am ₂ CO ₃ , filtered:—

EXAM^{N.} OF PP. IN GROUP II.—Removed from the filter into a porcelain dish and boiled with KOH, filtered:—

Filtrate:— Acidified with HCl, a white milky	Pp . removed from filter by a glass rod into a porcelain dish and boile with strong HNO_3 as long as any red fumes came off, added dflute H_2SO and stirred well:—					
Abs. of Group II.B.	No pp.:— Abs. of Hg	Added to so (Pres. of (olution excess of AmOH: k	olue solution		
	and Pb.	No. pp. :— Abs. of Bi.	Acidified the blue sola with with H ₂ S; filtered off the and boiled it with dilute Pp.: dissolved in a little boiling dilute HNO ₂ ,	e black pp. rapidly		
*	,		added AmOH in excess, then excess of HA, then K4FeCy6, chocolate red pp.:— Pres. of Cu.	H ₂ S, no pp. :— Abs. of Cd.		

EXAM^{N.} OF PP. IN GROUP III.A.

Dissolved in a little boiling dilute HCl, added pure NaOH in excess, boiled and filtered:—

Filtrate: added AmCl in excess, no pp.:— Abs. of Al.	Pp.: dried and fused on platinum-foil with Na ₂ CO ₃ and KNO ₃ , boiled the colourless (Abs. of Ct) mass when cold with water; decanted from the undissolved residue:—		
	Residue in the dish, dissolved by boiling with a little HCl, added several drops of KCNS: a blood-red coloration:— Pres. of Fe.	Soln. being colourless indi- cated abs. of Cr. Acidified with HĀ, boiled, and added PbĀ2, no. yellow pp.:— Abs. of Cr.	

EXAM^N. OF PP. IN GROUP III.B.

Rinsed the pp. off the filter with some cold dilute HCl, stirred well and filtered :—

contain no Ni fused	Filtrate: beiled until it no longer smelt of H ₂ S, added a crystal of KClO ₂ , boiled until the smell of Cl ceased, cooled and added pure NaOH in excess:—
a portion into a clear borax bead bead blue in both flames	Passed H ₂ S into the solution opp.
Rigit of Co	Abs. of Zn.

pp. in a few drops of HCl upon a watch-glass, dipped a loop of Pt-wire into the solⁿ and held it in the Bunsen-flame: a crimson colour was imparted to the flame, which appeared intense red through the indigo-prism: Pres. of Sr.

This coloration was followed by a yellowish green very persistent col^n : Pres. of Ba. Confirmed by spectroscope.

2. $Exam^n$ in the Wet way. Dissolved the rest of the pp. in as little boiling $H\bar{A}$ as possible; to a small part of the solution, when perfectly cold, added $CaSO_4$ -soln, an immediate pp. formed: Pres. of Ba.

To the remainder of the HĀ-solⁿ, proved to be acid to litmus, added K₂CrO₄ until the liquid appeared yellow, warmed and poured through a double filter:—

Pp., which contained all the Ba present, was rejected.	To the clear filtrate, which was orange-red in colour, added AmOH until the colour changed to light yellow, then added Am ₂ CO ₃ in excess and filtered:—		
	Filtrate was rejected.	Pp.: dissolved in as little boiling HA as possible; added to a small portion of the soln. CaSO ₄ -soln. and boiled: a pp. formed:—Fres. of Sr.	
		To the remainder of the $H\bar{A}$ -solution added H_2SO_4 , boiled, filtered, and to the filtrate, which gave no pp. after being boiled with more H_2SO_4 , added AmOH in excess and $Am_2C_2O_4$: no pp.:—	
		Abs. of Ca.	

EXAM^{N.} OF FILTRATE FROM GENERAL TABLE FOR GROUP V.

Evaporated the filtrate to dryness in a porcelain dish, scraped out the residue upon platinum-foil, and ignited strongly until white fumes ceased to appear: dissolved the residue off the foil by boiling with water to which several drops of HCl had been added, divided the solⁿ into two unequal parts:—

To the larger portion added several drops of H_8SO_4 and boiled, then AmOH in excess and several drops of $Am_2C_0A_4$; on heating, a slight pp. formed; filtered, and added to the clear filtrate Na_2HPO_4 , a white crystalline pp.:—

Pres. of Mg.

Into the smaller portion dipped a loop of platinum wire, and held it in the Bunsen-flame, an intense yellow coln.:—

Pres. of Na,

appearing crimson through the indigoprism:—

Pres. of K.

Confirmed the pres. of K by stirring a fresh portion of the soln. with H_2PtCl_6 on a watchiglass, a yellow pp. formed.

EXAM^{N.} FOR ACID-RADICLES.

The following acid-radicles have already been detected: (CO_3) , (NO_3) , (CIO_3) .

By reference to the Table of Solubility (1266), under the columns corresponding to the above metals, the only salt insoluble in acids is found to be $BaSO_4$; since Ba is present, and the substance is entirely soluble in acids, (SO_4) is absent.

The following acid-radicles have also been proved to be absent in the general examination for metals:—

(CrO₄), by the HCl-solution not becoming green on passing H₂S. (AsO₃) and (AsO₄), by no pp. forming in Group II.B.

(SiO₃) and (SiF₆), by no residue insol. in HCl remaining on evaporating to dryness with HCl for Group III.A.

(PO₄), by testing with AmHMoO₄ in the HCl solⁿ for Group III.A.

The organic acid-radicles (\bar{A}) and (\bar{T}) are absent, since there was no smell of burning on heating the substance.

The only commonly occurring acid-radicles remaining to be specially tested for are therefore Cl, Br, I, (C₂O₄), and (BO₃); and of these Br and I are probably absent, since no violet fumes of I or brown fumes of Br were evolved with strong H₂SO₄.

Boiled a portion of the substance with pure Na₂CO₃-solⁿ and filtered; acidified portions of the filtrate with—

-	HNO ₃	HĀ •
,	Added AgNO ₃ , a perfectly white pp., easily sol. in AmOH:— Pres. of Chloride.	Added CaSO ₄ , no pp. :— Abs. of (C_2O_4) .

Moistened a piece of turmeric-paper with the HCl-solⁿ. of the substance and dried it at 100° : no reddish-brown stain was produced:—Abs. of (BQ_3) .

Acid-radicles found: CO3, NO3, ClO3, Cl.

Metablic radicles: Cu, Fe, Co, Ba, Sr, Mg, Na, K, NH₄. Found Acid radicles: CO₂. NO₂, ClO₃, Cl.

PART IV

SECTION VIII

THE LABORATORY AND ITS FITTINGS; APPARATUS, CHEMICALS, AND REAGENTS

In Part IV directions are given which have been found by experience to be of value in fitting and furnishing a laboratory and in maintaining it in working order.

THE LABORATORY BUILDING, AND ITS FITTINGS.

1360. The methods of building, fitting, heating, and ventilating a Chemical Laboratory are of prime importance in securing the comfort, health, and convenience of the laboratory students.

The suggestions given in the following paragraphs are by no means exhaustive and they apply mainly to the preparation of a

chemical laboratory for analytical purposes.

It is evident that the details of furnishing and fitting must be influenced by the size, purpose and position of the laboratory. Provision must be made for the adequate lighting of the working-benches, for draught, waste and drainage; and these matters should be carefully planned before the laboratory building is erected, since much trouble and inconvenience may be caused by first building the laboratory and afterwards endeavouring to adapt the general fittings to an unsuitably designed structure.

The Building should provide for adequate air space and ventilation. The window-lighting should be from both the top and the

sides of the room, and on as ample a scale as possible.

The inner surface of the walls should either be of white glazed brick, which can be easily cleansed when necessary, or the brick or plaster surface should be covered with white lime-wash or distemper, which can be frequently and cheaply renewed. The windows should be glazed with colourless glass. This absence of solour in the walls and windows is necessary to enable the accurate judgment of colours to be made which is necessary to the analyst.

1361. Draught Arrangements.—The means of producing a draught for the removal of acid vapours and fumes from the chemical laboratory must be ample, and this is best secured by means of

an electric fan. As a rule stoneware draught-flues are preferable to those made of zinc or lead. These draught arrangements may be made to aid in the ventilation of the laboratory as well as to serve for efficiently exhausting the draught-chambers and bench flues.

Draught Chambers.—Around the walls, in the spaces between the windows, there should be large and small recesses, each of which is lined with white glazed tiles, has a glazed hood above, and is closed with a sliding sash in front; it should be provided with two outlets, one at the top and the other near the bottom, through which a strong draught is produced by the fan.

These draught-chambers should be provided with several gassupplies for Bunsen-burners, the taps being on the outside of the chamber; they should also be provided with means for lighting the interior. The bottom of the chamber may be covered with sheet-lead, and should have a small grid through which liquids escape into the waste. Some at least of these chambers should also be provided with a water-supply pipe.

A strip of rubber or fibre draught-excluder fixed along the top of the inner frame and lightly touching the glass prevents the escape of fumes when the sash is partly opened.

One or more of the draught-chambers should be reserved for the hydrogen sulphide apparatus (1371).

It will be found well to provide this H_2S -chamber with small doors, which are only opened when the gas is to be used: this prevents the gas from escaping in large quantity into the laboratory, as it would do if the large sash were thrown up for each operation. A frame containing pairs of little hinged glazed doors, each about six inches by eight, may be introduced beneath the sliding sash. The arrangement enables a student to open as much of the chamber-front as is necessary for passing the gas through a solution: while the whole sash can be raised for cleansing or replenishing the hydrogen sulphide apparatus.

One of the draught recesses is kept open for the steam-ovens, and the small boiler and condenser which are described in paragraph 1377.

1362. Dramage and Waste.—It is essential that the waste-service should be adequate. Lead pipes are unsuitable for this purpose, since they are liable to be corroded by acids, and are difficult to repair and to clear if they should become obstructed. The open twoigh system, made of stoneware, or of wood lined with sheet lead protected by a coating of bitumen, is to be preferred; the distinger from the penches and draught chambers should also

the drain may be readily examined and cleared by the removal of suitably arranged floor-boards.

Washing Sinks.—Two or more deep stoneware-sinks should be provided for general use, and should have large water-taps fixed above them at some distance from the back of the sink.

It will be found convenient if each sink is supplied with three taps. Two of these deliver hot and cold low-pressure water; and a third is connected with a high-pressure water service, upon which a fire-hose can be at once adapted. A stout wooden board pierced with small holes may be laid upon the bottom of the sink to serve as a strainer and to prevent solid fragments from getting into the outflow pipe; the soft wooden surface also tends to prevent the breakage of glass or porcelain vessels.

Note.—Lead-covered sinks lessen the risk of breakage arising from the dropping of apparatus while it is being washed, but they are easily corroded by acids and are less cleanly than stoneware. The wooden grid already mentioned, which is laid on the bottom of a stoneware sink, is a very efficient and permanent grid, and saves breakages.

1363. Working Benches.—In Fig. 66 facing page 331 a block of four laboratory benches is shown. Two of these are presented toothe observer, and are therefore visible in detail. The arrangement of four benches in one block is convenient, since it enables the washing-sinks and the water-taps to be placed at the end of each bench and yet to be accessible to each student. The surface of the bench is thus prevented from being wetted and soiled when apparatus is washed.

The Bench Sinks may be of stoneware, with perforated movable wooden false bottoms, which prevent breakage and serve as grids (Note, 1362). Half-inch outlet pipes unprotected by metal grids discharge into stoneware pots, the overflow from which into the drain-pipes is through a curved stoneware tube near the top. Any fine solid matter, which passes down the sink-pipe, settles in this trap-pot, and is got rid of by taking out the pot and emptying it into a pail at frequent intervals.

Any stoppage in the straight sink-pipe is readily pushed through with a stout cane. Over each sink there are three low-pressure water-taps for washing and for supplying condensers, and two high-pressure water-taps for aspirators.

A Stoneware Pocket is hung upon the end of each working bench to receive waste filter-papers, broken glass, and other solid refuse. This pocket is frequently emptied.

Bench Draughts.—Each bench is provided with an aperture leading to a strong draught, and into this aperture a light copper to porcelain hood can be fitted, which serves to carry away acid

vapours or badly-smelling gases. The aperture is closed with a wooden plug when the hood is not in use, so as not to diminish the draught on other benches.

The copper hoods are occasionally cleansed and painted with Japan-black, in order to protect them against corrosion by acid fumes: fixed light earthenware hoods are fragile, but present the advantage of not being corroded by acids. A glazed draughthood extending along the top of the centre shelf-rack of the bench has been found to be satisfactory: all heating and evaporation of liquids is then carried out on a raised shelf under this hood.

The Laboratory Bench may be made of stout teak or of American walnut, the wood of the front being stopped and varnished, and the working surface of the bench being saturated with oil and then well rubbed. The working surface is preserved by being occasionally thoroughly washed, dried, and rubbed down with a mixture of equal measures of boiled linseed oil and turpentine.

The bench may be 3 feet in height with a depth of 25 inches, and a space of 7 inches between the reagent-shelves above the bench will be found suitable for ordinary bottles. A gangway at least six feet in width should be provided between the rows of benches, and beneath this gangway the main drainage-troughs should run.

Two gas-taps are fixed on each bench for supplying Bunsen-burners. If gas is not available, Bunsen-burners for consuming methylated spirit or petroleum (1366) may be used with advantage. One electric glow-lamp or gas-burner is provided at a convenient height for illuminating the bench at night. If gas is used an incandescent mantle gas-burner is most suitable for this purpose since it gives the nearest approach to white light.

Bench Lockers.—Each working-bench is supplied with two lockers, and each locker has two drawers above it. In order to lessen the number of locks, which are liable to rust in the laboratory atmosphere, a flat iron bar which swings on a pin is fastened upon a staple by a padlock in front of the two drawers and locker. By this arrangement of lockers two students are enabled to use the same bench at different times.

Each locker should have a different key, but all the locks should be under the control of master-keys kept by the attendant and the teachers. The drawers should be about 3½ inches in depth, and may be suitably divided by wooden strips. A shelf in the locker should not extend more than half across, so as to leave standing toom for tall apparatus.

1364 Preparation and Gombustion Benches. Besides the ordinary working benches, there should be at least one long bench,

well supplied with gas and water-taps and with overflow-pipes. This bench serves for making up solutions, for distillations and for other large operations.

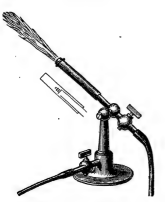
There should also be a long bench made of slate, and covered at a height of several feet with a metal hood, under which is a good draught-outlet. This bench must be supplied with a half-inch gas-pipe and taps, and with water-taps and waste-overflows: it serves to support water-baths, air-baths, combustion-furnaces, and gas furnaces of all kinds.

1365. Blowpipe-table and Blowpipes.—A table with a raised edge, and covered with sheet lead or zinc, is furnished with gastaps to supply standing gas-blowpipes, and luminous-burners for bending glass, and two or more Fletcher-bellows should stand beneath it. It is also well to have an electric blower near at hand to supply an air-blast to the blowpipes for lengthened heating operations.

The Fletcher Bunsen-blowpipe (Fig. 10, 12) will naturally find its place on this table. It is also well to keep a Bunsen gas-blowpipe (Fig. 67) on the table for more elaborate glass-blowing, and







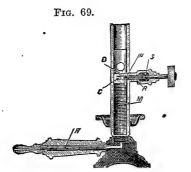
BUNSEN GAS-BLOWPIPE.



HERAPATH GAS-BLOWPIPE.

- a Herapath-blowpipe (Fig. 68) for smaller heating operations. The upper part of the Herapath-blowpipe can also be purchased without the foot, and this can be fitted into a Bunsen-burner and used on the working-bench.
- 1366. Spirit and Petroleum Burners and Blowpipes.— Heating and blowpipe-work must occasionally be done without a supply of coal-gas and without an air-blast. Recourse may then

be had to the Bunsen-burners and blowpipes which burn the vapour of methylated spirit or petroleum (Figs. 69, 70, 71).







PETROLEUM-VAPOUR BUNSEN.

In these forms of apparatus the burner is first heated by burning some methylated spirit in a circular gallery surrounding the burner,

Fig. 71.

BENZOLINE BLAST-LAMP.

and the supply of liquid coming into the base of the burner is then vaporised by contact with the hot burner.

In the case of the spirit-vapour Bunsen (Fig. 69) the liquid is supplied to the burner by gravitation from a vessel at higher level.

In the petroleum-Bunsen (Fig. 70) the petroleum is raised by the pressure of air which is forced by a little pump, shown on the right of the figure, into the upper part of the reservoir.

The blast of the blowpipe-lamp (Fig. 71) is maintained by the pressure of the vaporised liquid inside the reservoir.

- 1367. Chemical Store-room and Balance-room.—Opening into the general laboratory there should be five rooms, since it is usual to arrange a separate room for each of the following purposes—
 - (1) Balances.
 - (2) Chemical stores.

- (4) Physical apparatus.
- (5) Reference library and writing.
- (3) Gas and water analysis.

If space is limited, and only two rooms can be provided, one of the two is fitted with shelves for stores of chemicals and

apparatus. The other may be used as a reference-library: also for delicate metal apparatus, such as chemical balances, micro-

scopes, spectroscopes, and polariscopes.

The balances and delicate apparatus should stand on a broad shelf which is at a convenient height for sitting, the shelf being supported on iron brackets from the wall so as to secure the chemical balances against disturbance from the vibrations of the floor. The spectroscope and polariscope should have draw-curtains of black lining material, so arranged that light from outside may be excluded when the instruments are in use.

A large central table should be provided in this room for use in writing and reading.

A low table with a raised edge is also necessary for gas-analysis apparatus and for other experiments involving the use of mercury; and in order to provide against the loss of mercury spilt from the table, it will be well to cover the floor with linoleum with turned-up edges.

BENCH-APPARATUS.

Betailed descriptions of this apparatus may be obtained from the illustrated and descriptive price lists furnished by apparatus-sellers.

1368. A list of the apparatus which should be kept in each benchlocker will be found preceding page 1. The separate items of that list are repeated here, with the addition of details of dimensions and of other useful information.

In Part I. of the book, directions are given for preparing and fitting this apparatus, and the paragraph numbers in the lists below refer to these descriptions.

The Bunsen-burner with its rubber tube may be left attached to the gas-tap on the bench; all the other apparatus should be locked up in the bench-locker when it is not in use. Most of this apparatus is seen on the bench in Fig. 66, facing page 331.

The approximate decimal and English equivalents of length

and capacity are given to save reference or calculation.

1 Bunsen-burner (2), about $5\frac{1}{2}$ inches (14 cm.) in height, with $\frac{3}{6}$ -inch (1 cm.) tube, and with means of closing the air-holes.

1 Rose-top to fit the burner (2).

- 1 Fletcher's Argand-Bunsen burner (3), \(\frac{3}{4}\)-inch (1.8 cm.) size, is convenient for boiling the wash-bottle, but is not essential.
- 1 Piece of red or black rubber tubing, $\frac{5}{16}$ -inch (7.5 mm.) in internal diameter and 20 inches (50 cm.) long, to supply gas to the burner.
- 1 Test-tube stand with twelve holes, two of which are at least 1 inch (2.5 cm.) across; best without draining-pegs.

1 Test-tube brush (19).

12 Test-tubes, 5 inches (12.5 cm.) long, \(\frac{5}{8}\)-inch (1.5 cm.) in internal diameter.

- 2 Boiling-tubes, 6 inches (15 cm.) long, 1 inch (2.5 cm.) in internal diameter.
- 2 Round glass plates, ground on one side, 3 inches (7.5 cm.) across.
- 2 Porcelain evaporating-dishes with spouts, glazed inside and out, and 3 inches (7.5 cm.) in diameter.
- 2 Watch-glasses, 2 inches (5 cm.) across.
- 1 Corrical flask (Fig. 21.B., 36) of 4 ounces (120 c.c.) capacity.
- 1 Wedgwood-mortar, 4 inches (10 cm.) across, and pestle with wooden handle. 1 Galvanised iron tripod-stand, 7 inches (17.5 cm.) in height, with round
- 1 Galvanised iron tripod-stand, 7 inches (17.5 cm.) in height, with round top 4 inches (10 cm.) across (Fig. 21.B., 36); these dimensions are suitable to the above Bunsen-burner.
- 1 Square of coarse iron wire-gauze, 5 inches (12.5 cm.) in the side (Fig. 21.s., 36); best with the corners clipped off.
 - (A square of asbestos millboard may be used instead of the gauze.)
- 3 Plain glass funnels (Fig. 31, 43), two of them $2\frac{1}{2}$ inches (6.5 cm.) across, and one 2 inches (5 cm.) across.
- 3 Beakers without lips, wide form, of 3, 4 and 6 oz. (100, 120 and 200 c.c.) capacity.
- 3 Glass rods, round at the ends, 7, 6, and 3 inches (17.5, 15 and 7.5 cm.) in length (15).
- 1 Piece of platinum foil, 1 inch square (2.5 cm. square).
- 2 Pieces of mounted platinum wire (14), each 2 inches (5 cm.) long and about as stout as an ordinary sewing needle.
- 1 Blowpipe, Black's japanned tin (Fig. 9, II).
- 1 Pipe-clay triangle, 2 inches (5 cm.) along its side (Fig. 65, 1308).
- 1 Wash-bottle (17), made by fitting a conical 18-ounce (500 c.c.) flask, which is at least 1 inch (2.5 cm.) across the inside of the neck.
- 1 Wooden filter-stand (Fig. 33, 43), rod 12 inches (30 cm.) high, two rings on one boss, each $2\frac{1}{4}$ inches (5.5 cm.) across, foot 5 by 8 inches by $\frac{1}{2}$ inch (12.5 by 20 by 1.3 cm.).
- 1 Pair of polished brass crucible-tongs, 6 inches (15 cm.) long.
- 1 Small horn spatula, 3½ inches (9 cm.) long.
- 1 Wicker oval draining-basket, 10 inches by 8, and 4 inches deep. Cut filter-papers, $4\frac{1}{2}$, $3\frac{1}{2}$, and $2\frac{3}{4}$ inches (11.5, 9 and 7 cm.) across.

APPARATUS FOR GENERAL USE IN ANALYSIS.

1369. The Following Apparatus should be kept in the laboratory for the general use of students who are doing chemical analysis.

One set will suffice for about twelve students. Each article should have its own place in the laboratory, and should be replaced immediately after use, since it may be required by other students.

- 3 Nests of three or four of the smallest brass cork-borers.
- 6 Triangular files without wooden handles.
- 3 Thin round files without wooden handles.
 - Several lengths and pieces of hard glass tubing, about \(\frac{1}{8} \) inch (3 mm.) internal diameter.
 - Ignition-tubes, 3 inches long by \(\frac{1}{4} \) inch across (7.5 by 0.6 cm.).
- Corks as free as possible from holes and cracks, sizes varying from \$ to \$\frac{3}{8}\$ inch across (1 cm. to 2 cm.).

Several retort-stands of galvanised iron (Fig. 34, 44), upright rod 16 inches (40 cm.), foot 6 by $3\frac{1}{2}$ inches (15 by 9 cm.) with three brass or gun-metal rings, diminishing from 3 inches (7.5 cm.) across, and metal clamp (Fig. 22, 35).

3 Tubulated retorts of 6 ounces (200 c.c.) capacity.

6 Thistle-funnels, of assorted length.

6 Clock-glasses, about 4 inches (10 cm.) across.

Pieces of red or black rubber tubing, $\frac{3}{16}$ inch (5 mm.) in internal diameter and $1\frac{1}{2}$ inches (4 cm.) long.

A good supply of wooden spills, or thin strips of wood.

A good supply of wax tapers, about 11 inches (27 cm.) long.

- 1 Spirit-lamp, 4 ounces (120 c.c.) in capacity, with earthenware wick-holder and ground-glass cap (5).
- 1 Charcoal borer of conical form.
- 4 Tin filter-dryers (Fig. 39, 48).
- 4 Porcelain crucibles, 11 inches (4 cm.) across, with covers.
- 1 Iron mortar, 8 inches (20 cm.) across, with pestle.
- 1 Fletcher's No. 5 foot-bellows (Fig. 11, 12), and a blowpipe-table covered with sheet zinc or lead.
- 1 Fletcher's Bunsen-burner blowpipe, with central blast, and two taps (Fig. 10, 12).
- 1 Bunsen-blowpipe, and one Herapath-blowpipe (1365).
- 1 Fletcher's solid-flame burner, with tripod for large dishes (Fig. 2, 3).
- 1 Fletcher's injector-furnace for strongly heating large crucibles.
- 1 Microscope, with at least a 4-inch (6 mm.) object glass.
- 1 Spectroscope, Bunsen's table form (65).
- 1 Spectroscope, direct vision, pocket form (64).
- 1 Indigo-prism, stoppered, and filled as is described below:-

Clear indigo-solution is made by mixing commercial sulphindigotic acid with about ten times its measure of strong sulphuric acid, and letting this settle for several days.

The prism is nearly filled with this solution of indigo; the dry stopper is inserted into the dry neck and is securely fastened down with fine copper wire; melted paraffin wax is then allowed to flow over the junction of the clean dry stopper and neck, and to harden over it.

1 Agate mortar 3 inches (7.5 cm.) across, and pestle.

This mortar is used for reducing hard substances to fine powder, after they have been already broken into minute fragments. The process of powdering in the agate mortar must be effected by trituration, since if blows are struck with the pestle the mortar may be broken.

2 Small leaden cups, about 1½ inches (4 cm.) across and ¾ inch (2 cm.) deep

(654). These are made by beating sheet-lead into shape. 2 Copper water-baths, 5 inches (12-5 cm.) across, and with four diminishing

rings (Fig. 27, 38).

One large bath with several openings (Fig. 28, 38) may with advantage replace these smaller baths. The baths should be about two-thirds filled with water, and the flame should be so adjusted that the water is kept just boiling. Evaporating basins placed on the bath cannot be heated above 100° C.

- 2 Pairs of conical flasks, fitted as is shown in Fig. 53 (176).
- 2 Tubulated flasks, 4 ounces (120 c.c.) in capacity (Fig. 56, 474).
- 2 Clarke's retorts with condensers (Fig. 57, 542).

A box of assorted pieces of wood charcoal, as free as possible from cracks and from bark.

Hydrogen Sulphide Apparatus.

1370. Hydrogen sulphide Gas is constantly required by the analyst, and should be readily obtainable at any time in a constant stream; but arrangements must be made for preventing this poisonous and badly-smelling gas from polluting the atmosphere of the laboratory. These objects, together with economy of materials, are better secured by the use of a large apparatus in common by many students than by the employment of a small private apparatus by each student.

The gas is most readily obtained by the action of somewhat diluted cold commercial hydrochloric acid upon fragments of ferrous

sulphide.

1371. General Hydrogen sulphide Apparatus.—The apparatus in which the gas is prepared has assumed many forms. The Kipp-apparatus (Fig. 73, 1272) is probably in most general use, but the apparatus shown in section in Fig. 72 and described below is also recommended by its simplicity, efficiency, and cheapness.

Two large tubulated bottles, A and B, of at least 2 quarts ($2\frac{1}{2}$ litres) capacity each, are fitted as is shown in the figure. The tubulures of the bottles are connected together by a piece of broad rubber tubing, and the bottle B is also connected with the wash-bottle C by narrow rubber tubing. Rubber stoppers should be used, since they are more durable and gas-tight than ordinary corks. The delivery tube D is connected with the doubly-bent tube from the washing-bottle by means of a rubber joint, which can be partially or entirely closed by means of a screw-clamp E.

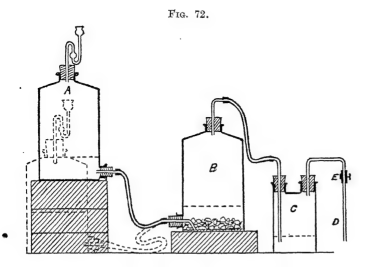
In charging the apparatus, the bottle B is laid on its side and is then filled with lumps of ferrous sulphide, which should be free from powder, since the powder is apt to choke the rubber connecting-tube, and to continually generate gas in B when the production should be arrested. The washing-bottle C is then partly filled with water, and the three bottles are connected. The clamp E is now closed, and the bottle A is about half filled with a mixture in equal measures of strong commercial hydrochloric acid and water.

When the gas is required, the bottle A is raised upon a suitable stand, and the clamp E is gradually opened until a regular stream of bubbles passes through the water in C.

A part of the acid will flow from the bottle A into the bottle B, and will cause hydrogen sulphide to be evolved from the ferrous sulphide; and the gas will be forced out of B by the pressure of the acid which remains in A. Before it escapes through the tube D, the gas is freed from small drops of liquid, containing HCl and

FeCl₂, by passing through the water in C. The levels of the liquids in the bottles A, B and C, while the gas is being produced; are shown by the horizontal dotted lines.

, When the gas is no longer required, the screw-clamp at E is



LARGE HYDROGEN SULPHIDE APPARATUS.

gradually tightened until the stream of bubbles through the water in C is arrested. The gas then accumulates in B, and forces the acid out of B back into A. If the glass tube at the bottom of B is bent, as is shown in the figure, the acid will be almost completely expelled. This will cause the production of the gas to cease, unless FeS-powder rests on the bottom of the bottle B. The evolution of gas will recommence, however, as soon as the clamp E is opened. During the hours of laboratory work A should remain raised on its stand, and the apparatus will then always be ready for use. At other times A should be lowered into the position represented in dotted outline in the figure. B should be permanently somewhat raised in order to facilitate the outflow of the acid when A is lowered.

The escape of the hydrogen sulphide gas from solution in the acid is much lessened in amount if a funnel containing water or glycerine is fitted into the stopper at the top of A, or if a loosely-fitting glass stopper is dropped into the neck of the bottle.

· When the acid is spent and no longer causes evolution of gas from the FeS, it is poured away and replaced by fresh acid. Lumps of FeS are also occasionally placed in B, as the substance becomes dissolved.



THE KIPP APPARATUS.

1372. The Kipp-apparatus (Fig. 73) may also be used for generating hydrogen sulphide. It is similar in its action to the one already described, but the acid vessel (f) is placed above the generating vessel (c), and the two are connected together rigidly when the apparatus is fitted up for use.

In charging this apparatus the tube of the upper vessel (f) is withdrawn from the ground neck of the middle vessel (c), and lumps of FeS are introduced into e through the opening while the apparatus is laid upon its side. The dry clean neck is then greased, and the tube of (f) fitted into the neck while the apparatus is still lying on its side. The apparatus is then placed erect, the bottom vessel (a) is stoppered, and the delivery-tube with closed stop-cock is fitted in at (d). The diluted HCl is now poured in through the neck of (f), and the diffusion of H_2S from the acid checked by fitting in a water-valve as shown. When the acid is spent it is withdrawn by removing the stopper in the bottom vessel (a).

The hydrogen sulphide apparatus should stand in a draught-chamber, which is lined with glazed white tiles and is furnished with a good draught. The bottom of the chamber should be covered with sheet-lead, furnished with a grid and drain-pipe for the outflow of the very foully-smelling spent acid, or this liquid may be poured away outside the laboratory building. The chamber must not be used for other purposes, and must be suitably lighted at night.

A large sash or door must be opened when the spent acid is poured away and the apparatus is recharged within the chamber; but a small door only should be used for introducing the solutions through which the gas has to be passed. If the front of the chamber is opened by means of a glazed sliding-sash, a frame containing pairs of glazed hinged doors, each about six inches by eight, may be introduced into the grooves beneath the sash. These doors enable the apparatus to be used with the least possible opening of the closet.

Such an apparatus should serve for the use of twelve or more students, according to the character of the work which is being done.

1373. An Apparatus for Private Use may be fitted as is shown in Fig. 74. The larger bottle contains pieces of ferrous sulphide, upon which HCl, diluted with an equal measure of water, is poured through the thistle-funnel. The gas passes through a little water in the smaller bottle, and thence into the solution to be saturated. The use of a small apparatus by each student in a large laboratory is not to be recommended, since it causes unnecessary waste of time and materials, and does not admit of satisfactory control.

1374. Passing Hydrogen sulphide.—Each student should

keep a glass tube D (Figs. 72, 73, 74), which may be fitted upon the hydrogen sulphide apparatus in order to conduct the

gas into a solution.

When a liquid is to be saturated with gas, the end of this tube should be passed down to the bottom of the vessel which contains the liquid. A regular stream of gas is then caused to bubble through the liquid by gradually opening the clamp E (Fig. 72), or the stopcock d (Fig. 73) of the larger apparatus, or by introducing a sufficient quantity of acid into the larger bottle of the small apparatus (Fig. 74).



MALL HYDROGEN SUL-PHIDE APPARATUS.

In order to avoid unnecessary waste of gas, the liquid should be of sufficient depth to absorb much of the gas, and the bubbles should not pass too fast to be readily counted.

The passage of the gas should be continued for four or five minutes if the quantity of metal to be precipitated is large, and the liquid must smell strongly of the gas after the liquid has been shaken, and the air above it has been blown out. The tube D must be thoroughly cleansed immediately after use.

Liquid Hydrogen sulphide can often be purchased in metal bottles. When the valve of the bottle is slightly opened, a stream of the pure gas will escape. A store of ready-made hydrogen sulphide may be kept in this form, and the trouble of maintaining the apparatus for preparing the gas may thus be obviated.

1375. Hydrogen sulphide Solution.—It will be found convenient to keep a bottle of distilled water saturated with hydrogen sulphide. This solution is prepared by passing a stream of the gas through water which nearly fills the bottle (Fig. 81, 1395). On removing the bottle, closing its mouth tightly with the thumb and shaking its contents violently, no suction must be felt on the thumb, but a slight outward pressure: this proves that the water is saturated with the gas.

DISTILLATION OF WATER.

1376. As has been already stated (31), water ordinarily contains certain solid substances in solution, which render it in a chemical

sense impure. Such water is therefore unfit to be employed for the processes of solution and of washing precipitates, since any impurity thus introduced during the analysis would be detected and considered to have been present in the original substance.

The quantity of these impurities which is present in any particular water-supply will depend upon the nature of the soil with which the water has been in contact before its collection. The water-supply in some districts will be found to be almost perfectly pure, when it is subjected to the tests for impurity (31).

Rain-water which has fallen through pure air, and has been carefully collected, will yield little or no indication of dissolved impurity. The water-supply to the majority of laboratories will, however, be found to be unfit for analytical purposes, until it has been freed from dissolved solid substances by being subjected to the process of distillation.

In most towns distilled water can be purchased; but it is preferable to distil all the water required for use, either in the laboratory itself or in its immediate neighbourhood. For this purpose steam is generated and condensed by a suitable cooling arrangement.

The steam may be obtained from a steam-boiler which is in use for other purposes, or from a special copper boiler heated by solid or oil fuel or by a gas-burner. It is advantageous and economical to utilise the steam for heating the steam-ovens before it passes to the condenser: and water-stills are now made in all sizes, which effect these purposes satisfactorily.

The apparatus described in par. 1377 is now in general use in students' laboratories and is recommended both for convenience and for economical working.

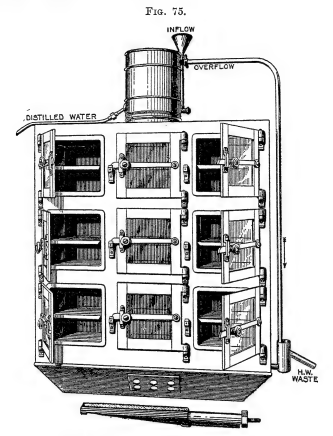
Smaller and simpler stills of a similar kind are obtainable when they are required for individual use only.

The hot outflow of condensing water from the still may be made to pass through a collecting-tank on its way to the waste service, and hot tap-water may then be drawn from this tank when required.

1377. The Distillation of Water may be combined with the Heating of Steam-ovens and Steam-bath by the following apparatus, known as the Brown-still, which may be conveniently placed in one of the open draught-recesses in the laboratory-wall.

A front view of this still is shown in Fig. 75; and it will be seen that a shallow boiler, forming the bottom of a range of nine steam-ovens, is heated by a long Fletcher gas-burner. The steam passes from the boiler around the ovens, and is condensed by a special and very efficient condenser which is mounted on the top

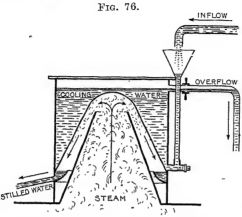
of the range of steam-ovens, the distilled water thus produced being taken away by a pipe to a stoneware storage vessel. The top of the casing may have different-sized holes cut for dishes, and closed by lids when not in use: these serve for water-baths and hot plate (Figs. 26, 27, 28, par. 38), but they lessen the yield of distilled water.



THE BROWN STEAM-OVENS AND STILL.

There is an inflow of cold water from the water service to the bottom of the condenser, and the hottest water flowing away from the top of the condenser passes to a small feed-cylinder attached to the side of the boiler; a part of this water serves to maintain the level of water within the boiler, and the rest flows away through a hot-water storage tank into the waste service. The sectional drawing (Fig. 41, 48) will explain this feed-arrangement.

The sectional drawing (Fig. 76) renders evident the construction of the condenser, which exposes the steam to a large conical dome



THE BROWN-CONDENSER (IN SECTION).

kept cold by the circulating water. The water formed by condensation in the dome is collected by an annular internal channel, which is connected with the outflow for the distilled water.

A detachable plate is screwed down over an opening in the boiler, and renders possible the removal of incrustation by scraping or by means of

strong commercial hydrochloric acid diluted with four times its volume of water.

The oven-doors should be made of sheet-glass framed in copper, so as to render the interior of the ovens visible when the doors are closed; and in order to obtain a temperature near 100° C. in the oven, the air-inlet pipe to the ovens should be surrounded by steam so as to deliver heated air into the bottom of the ovens. An air-outlet is provided at the top of each oven.

The whole apparatus is made in sheet copper, and the interior surfaces of the condenser-dome and of the outflow pipe for the distilled water are thickly tinned to prevent contamination of the distilled water with copper.

When the gas-burner and supply of condensing-water have been properly adjusted, the apparatus may be allowed to work continuously without requiring any further attention.

RECOVERY OF SILVER AND PLATINUM FROM THEIR RESIDUES.

1378. Waste scraps of platinum and silver, and solutions and precipitates which contain these metals, should not be thrown away, but should be preserved in two large jars which are specially

reserved for them, and are labelled Silver Residues and Platinum Residues respectively. Platinum chloride and silver nitrate may be obtained from these residues by the methods described below.

SILVER RESIDUES.

1379. Some HCl is poured into the jar which contains the silver residues, and the acid is thoroughly mixed with the contents of the jar. The precipitate is then allowed to subside, and, after the liquid has been decanted, this precipitate is thoroughly washed by decantation: it may then be treated either by paragraph 1380 or 1381.

1380. The wet precipitate is rinsed into a flat shallow dish, and the layer of water is acidified with H_2SO_4 . Strips of Zn, free from Pb, are then laid upon the silver precipitate, and the whole is allowed to stand quietly for several hours. The black spongy mass which remains is metallic silver: it is separated from any remaining Zn, and is washed until it is free from H_2SO_4 .

This metallic deposit is dissolved by heating it with pure strong HNO₃, which has been diluted with its own measure of water. The solution is evaporated to dryness; the solid residue is dissolved in water, and is once more evaporated to dryness in order to remove free acid completely. The residue is then dissolved in the proper proportion of water to give reagent 54 (1410). Any residue insoluble in HNO₃ is returned to the silver residue bottle.

r38r. The well-washed precipitate which has been obtained from the silver residue bottle, after the addition of HCl in excess (1379), is filtered off and dried. It is then mixed with twice its weight of a mixture in equal proportions of K₂CO₃ and Na₂CO₃. This mixture is transferred to a clay crucible which is heated in a furnace. After the contents have been kept in a fused condition for four or five minutes, the crucible is removed and its bottom is tapped several times on a brick so as to cause the melted globules of Ag to unite. When the crucible is cool it is broken up, and the button of Ag is removed and well washed. It is then converted into AgNO₂-solution, as is described in paragraph 1380.

PLATINUM RESIDUES.

1382. The liquid in the platinum residue bottle is shaken up with the precipitate, and the whole is evaporated to dryness in a porcelain dish. The residue is then heated strongly for some time. When the dish is cool, water is poured upon the residue and boiled with it, and the aqueous solution is then decanted off. Solution of

oxalic acid is now poured upon the residue in the dish and evaporated to dryness, and the residue is once more strongly ignited.

The residue of metallic platinum is thoroughly washed with boiling water, and is dissolved by heating it with HCl, to which one-third its measure of HNO₃ has been added. The solution is evaporated to dryness over a water-bath. HCl is then poured in, and the liquid is once more evaporated to dryness and heated for some time on the water-bath. This residue, when dissolved in water, forms reagent 43 (1410).

Waste scraps of platinum foil and wire should be carefully preserved. They are cleansed by boiling them with HNO₃ and washing them well with water. They are then dissolved in a mixture of HCl and HNO₃, and converted into platinum chloride solution as is described above.

CHEMICALS AND REAGENTS.

1383. In the Following Paragraphs general Directions are given for the preparation of the reagents and of the test-substances which are required in the preceding Analytical Course.

Tabulated lists of the names and formulæ of these substances follow the general directions. A reference number is attached to each substance, and certain necessary details concerning its preparation and the tests for its purity are also supplied.

The pure chemicals which are required in analysis can be purchased, and it will be found more economical as a rule to buy them than to prepare them.

The preparation of the solutions and the dilution of acids and other liquids should, however, always be performed in the laboratory, since this lessens the cost both of purchase and of carriage.

Substances suitable for analysis are suggested in paragraphs 1415-1419.

1384. Solution and Dilution.—Many reagents and test-substances must be dissolved or diluted before they are used. As these processes of solution and dilution must be constantly carried out in a laboratory, it is important that the methods employed should be as simple and rapid in execution as possible, in order that the expenditure of labour and of time may be minimised.

It is also of great importance that the solutions should be of appropriate strength. It is usually advisable that the same liquid should be of different strengths when it is used as a reagent and as a test-solution. In either case a strength can be selected which generally yields the most satisfactory results. If the solution is

stronger than this, chemicals are wasted; if it is weaker than this, the reaction may not be satisfactorily obtained. Hence it is advisable to keep these liquids in readiness for the student, rather than to leave him to prepare them doubtfully or wastefully when they are required.

And in order to lessen the time involved in preparing these solutions, it is well to make each solution in some quantity and to keep it in stock.

1385. Normal Solutions.—It is customary to prepare all solutions with a strength which is some multiple or submultiple of the strength of a "normal" solution; thus the strength may be three times normal, normal, half normal, one-tenth normal, and so on.

By a "normal" solution is meant a solution which contains one gram-equivalent per litre; thus one litre of a normal solution of HCl contains 36.46 grams of HCl, one litre of a normal solution of Na₂CO₃ contains $\frac{Na_2CO_3}{2} = 56$ grams per litre, since this weight is chemically equivalent to 36.46 grams of HCl.

The advantage of adopting this system is that the volume of one solution which should be used to react quantitatively with another solution is known. Thus 1 c.c. of any normal acid-solution will neutralise 1 c.c. of a normal alkali-solution. Similarly 1 c.c. of normal BaCl₂- solution will precipitate quantitatively the sulphate in 1 c.c. of normal H₂SO₄.

1386. The Apparatus required for Preparing Solutions includes—

A common balance, with earthenware pans, and the beam below.

A more delicate balance.

A set of brass weights from 1 kilogram to 1 gram.

A strong measuring-cylinder to deliver one litre, and with graduations for every 5 cubic centimetres (c.c.).

Several large wide-mouthed gallon (5 litre) jars, some of stoneware or earthen-ware, others of common green glass.

Several large funnels of glass or of porcelain.

A convenient store-bottle for liquids is a well-cleansed Winchester-quart, the bottle in which acids and other liquids are constantly supplied to the laboratory.

The description of the processes for preparing liquid reagents is divided into (1) Dilution of Liquids (1387), (2) the Solution of Solids (1388-1393), and (3) the Solution of Gases (1394-1400).

DILUTION OF LIQUIDS.

r387. In the Lists of Diluted Liquids which Follow, the proportions by measure, in which the liquids are to be mixed with distilled water, are stated. Liquids may be mixed in these proportions by measuring them before they are mixed. But the process is often simplified by measuring the height from the bottom of the cylindrical vessel in which the mixture is to be made, to the level which the mixture is to reach. This height is then divided in the requisite proportion; and the division is marked by a file or a diamond, by a painted line, or by an india-rubber ring. Each liquid is then poured in to its own level, and the liquids are thoroughly mixed by shaking the bottle. The mark, when once made, will always serve for diluting the same liquid in the same vessel.

This method serves for numbers 2, 3, 4, 6, and 7 (1409).

In the preparation of dilute $H_2SO_4(1, 1409)$ special precautions are necessary, because the dilution of the strong acid produces much heat, see Remark 1 (1409).

SOLUTION OF SOLIDS.

r388. In the Lists of Dissolved Solids which Follow, the weight of the solid to be dissolved in water, and then made up to one litre in preparing each solution, is placed in the Fifth Column.

The number of grams of the solid to be dissolved in a Winchester-quart of water may be obtained by multiplying this weight by 2-4; since an average Winchester-quart bottle, filled about an inch above its shoulder, holds 2,400 c.c., or approximately 2,400 grams, of distilled water.

In a similar manner the weight of water held by any other bottle or vessel may be obtained with sufficient accuracy for this purpose by measuring its capacity in c.c., and counting this number as the weight of water in grams.

Pure solid chemicals are usually sold in the crystalline condition. Hence it may be assumed that the crystalline substance is to be used whenever this is possible.

1389. If a Small Quantity of a Finely-powdered Solid has to be dissolved, or if the solid is extremely soluble in water, the solution is often most simply made by introducing the weighed solid at once into the water contained in a stoppered bottle or a beaker, and then shaking or stirring the water until solution is completed. This method is successful with No. 12 (1409), with Nos. 19, 23, 27, 30, 43, 44, 49, 51, 54, 58, (1410), and with many of the substances in paragraphs 1411, 1412.

1390. For Dissolving Large Quantities of Solids two methods are described below. The first (1391) is useful when the solution must be rapidly prepared, and is of general application to the solution of solids; but the second (1392) is decidedly preferable for many reasons, and should always be employed if possible.

FIRST METHOD.

1391. Weigh out the required quantity of the solid substance, and heat it with distilled water in a large flat-bottomed flask on a sand- or water-bath with occasional shaking, or in a large porcelain dish over a Bunsen-burner with a rose top or over a Fletcher-burner.

This solution would often crystallise as it cooled, unless it is further diluted with water; hence, if it is not clear, it should either be filtered in a hot filter-jacket (see *Note*), or it may be at once poured into some cold distilled water contained in a large beaker, which has been marked to show the quantity of liquid required to fill the store-bottle.

While the liquid is being poured out of the dish, a wet glass rod should be held vertically and pressed against the lip of the dish, and the bottom of the dish should touch the top of the beaker. These precautions will prevent the liquid from running down outside the beaker.

Note.—The Filter-jacket, shown in Fig. 77, is a double-walled vessel, which is shaped to fit the funnel, and has a hollow projection (a) communicating with the interior of the jacket. Before the apparatus is used, it is about half filled with water, and supported on a tripod-stand. The water is then heated to boiling by placing a Bunsen-flame beneath the projection (a), which is filled with water communicating with that in the jacket. In this way hot water will circulate throughout the copper jacket, and the funnel supported within may be kept hot during filtration.

The solution is now diluted with distilled water to the required volume, if this has not been done already: the liquids are mixed well by stirring; and if the solution is turbid, it is covered and allowed to stand until it is clear. It is then poured or siphoned off into the store-bottle, the sediment being left behind.



THE FILTER-JACKET.

If the liquid is wanted in haste, it may be filtered into the bottle. For this purpose a double filter or a fluted filter may be used. If the liquid is filtered before it has been diluted to the full extent, the quantity to be passed through the filter will be lessened and time will be economised.

The Operation of Filtration, as it is described for ordinary

analytical purposes in paragraph 43, is very slow when a large quantity of a solid has to be separated from a liquid. This is partly due to the fineness of the pores of the filter-paper, and partly to the fact that the paper is in close contact with the sides of the funnel.

Methods are described below which serve better for dealing with large quantities of material to be filtered, such as are used in the preparation of pure chemicals and of stock solutions. It will be noticed that the procedure to be employed varies with the character of the solid.

Gelatinous or Non-granular Solids are conveniently separated from the liquids which contain them, by filtration through a large surface of fine linen.

One method for effecting this is shown in Fig. 78. The linen is first washed in hot water until it is free from glaze: it is then placed in a large funnel and made to assume the shape of a bag, as is shown by the dotted line. The bag is kept in its place by pressing a ring of wood firmly into the top of the funnel, as is shown in shaded section. When the filtration is finished, the bag may be squeezed so as to remove the adherent liquid more completely from the solid.

A simple modification of this method consists in making a square wooden frame about six inches across, and driving in a wire nail at each of its corners. The heads of the nails are then cut off so as to form four pins upon which the linen can be hung in the form of a bag. The funnel is thus dispensed with.



LINEN FILTER.

Granular or Crystalline Solids may be conveniently separated from the liquids which contain them, by means of a circular perforated porcelain plate 2 or 3 inches (4 to 7 c.m.) in diameter.

The perforated plate is dropped into the funnel and remains about $1\frac{3}{4}$ inches (4 cm.) from the apex of its conical interior surface.

The plate is then covered with a disc of filter-paper or of linen, which should be very slightly larger than the plate. After the moistened filter has been pressed into close contact with the inside of the funnel, the liquid which is to be filtered is poured upon the filter.

Filtration will take place much more rapidly in this filter, than in an ordinary paper filter which is in con-

tact with the funnel by its whole aurface.

• Perforated porcelain funnels, known as Buchner funnels, satisfactorily replace the combined perforated plate and funnel.

THE SECOND METHOD.

r392. This depends upon the fact that the density of the solution becomes greater as the quantity of solid dissolved in it is increased. Hence if the solid is immersed just beneath the surface of the water, a circulation of the liquid is produced. The water which has been in contact with the solid, and has been increased in density by dissolving it, constantly sinks and is replaced by the comparatively

lighter liquid from below. This circulation of water over the solid substance gradually dissolves it without the use of heat, and the process requires no attention after it has once been started. The process may be carried out as is described in (a), (b), or (c) below.

- (a) The most simple method of applying this principle is to lay the weighed solid upon a piece of dry well-washed muslin. This is then tied up into a little bag, and is suspended in the upper part of the water contained in a jar of common stoneware or earthenware or of thick green glass. A glass rod placed across the mouth of the jar serves to hang the bag upon. The process of solution by this method is very rapid.
- (b) A glass cylinder A (Fig. 79) which is open at both ends (such as a broken beaker, flask, or bell-jar) has one end covered with muslin, which is fastened on by a rubber ring and is shown by the lower dotted line. This cylinder with a muslin bottom is supported in the jar by means of two pieces of glass Fig. 79.

rod or tube, bent as is shown at c c c.

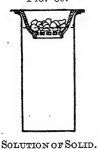
Distilled water is poured into the jar until it buries the muslin to the depth of about half an inch. The weighed substance is then placed in the cylinder, and the whole is allowed to stand until the solid has been dissolved. With large quantities of substance this will usually require several hours; with smaller quantities about twenty minutes or half an hour will be necessary.

(c) A convenient substitute for the glass cylinder with muslin bottom is an earthenware colander (Fig. 80). If the jar is too large, a stoppered bottle of suitable size may be placed in it so as to raise

SOLUTION OF SOLID.

the water level sufficiently to reach the substance.

Fig. 80.



The processes of solution and filtration may be performed simultaneously by laying a piece of filter-paper on the bottom of the colander, or upon the muslin before it is fastened on the cylinder. The presence of even the coarsest filter-paper will, however, considerably retard the process of solution.

Large quantities of substance are most conveniently dissolved by starting the process overnight; the solution will then be completed by the morning. In this case the retardation caused

by the filter-paper will be immaterial. The paper is, however, usually unnecessary, since any fine particles which have passed into the water will settle during the night, and the liquid can therefore be easily decanted from the sediment. It must be borne in mind that the volume of the water is increased by the solution in it of a salt, and especially by salts which contain water of crystallisation. Hence the quantity of water which is placed in the jar must be less than the volume of solution required, and the liquid must be made up to the required volume after the solution has been effected.

1393. Preparation of Saturated Solutions of Solids.—A solution is said to be saturated, when it contains the largest amount of the substance which the liquid can dissolve at the temperature of the solution. The quantity which can be dissolved usually varies widely with the temperature, and in the case of solid substances commonly increases with rise of temperature. Saturation is always understood to be effected at ordinary atmospheric temperature in the following text.

The methods described in paragraph 1392 are very convenient for the preparation of saturated solutions. The solution is known to be saturated when the solid supplied to the upper part of the liquid ceases to be any further dissolved.

In the case of a finely-powdered substance, which will remain long suspended in water, a saturated solution may be easily obtained by shaking the powder up with water. The powder is then allowed to subside, and the clear solution is decanted. The two following solutions may be prepared in this way.

1. Calcium sulphate Solution, No. 11 (1409), is made by filling a Winchester-quart up to its shoulder with distilled water, then pouring in some plaster-of-Paris powder, stoppering the bottle, and shaking it well. The liquid will remain milky for a long time; and since the water is thus kept in contact with a large surface of the solid it will become saturated with CaSO₄ before the powder has settled.

As soon as the liquid has become perfectly clear by settling, the solution is decanted into another bottle for use. The sediment remaining in the first bottle is again shaken up with a fresh quantity of water and the bottle is set aside. It will then furnish a fresh supply of the clear solution when necessary.

2. Lime-water, No. 36 (1410), is prepared by introducing freshly-slaked lime into a Winchester-quart bottle, filled to its shoulder with tap-water, and proceeding precisely as is described for the preparation of calcium sulphate solution in the preceding paragraph.

SOLUTION OF GASES.

1394. Many gases, which are more or less soluble in water, are most conveniently applied as reagents in the dissolved condition. Those solutions which are most frequently used, such as solution of ammonia, of hydrogen chloride and of sulphur dioxide, can be readily and cheaply purchased.

Of the remaining gaseous solutions: hydrogen sulphide solution should be kept in readiness, but the gas may be passed through the test liquid instead (1374). Nitrogen tetroxide solution is replaceable by a freshly-acidified solution of potassium or sodium nitrite. And as a substitute for chlorine-water, either brominewater, or freshly-acidified solution of bleaching-powder may be employed.

It will be seen, therefore, that the preparation of gaseous solutions is by no means indispensable.

1395. Preparation of Saturated Solutions of Gases.—All the solutions of gases (1396-1400) should be prepared in the open air or under a good draught, since the gases have an unpleasant smell and are injurious when inhaled.

• Each gas is made to pass from the generating apparatus down a delivery tube (a, b, Fig. 81) to the bottom of some cold distilled water, contained in a bottle which can be closed by a well-fitting stopper. The water is kept cool, if necessary, by immersing the bottle in a pan of cold water. Bubbles of gas Fig. 81.

are thus made to pass in a rapid stream through the whole height of the liquid, and a portion at least of each bubble is dissolved.

The solution is saturated as soon as the gas is not further dissolved. This may be judged in some cases by observing whether the bubbles cease to diminish in size during their ascent through the liquid. But a better test of saturation consists in withdrawing the delivery-tube, at once closing the bottle with the stopper or the Saturation of thumb, and shaking it well. If pressure is felt



WATER WITH GAS.

from within, due to the liberation of some of the dissolved gas by the agitation, the solution is saturated. But if suction from within is felt, due to the solution of the gas filling the upper part of the bottle, the liquid is not yet saturated with the gas.

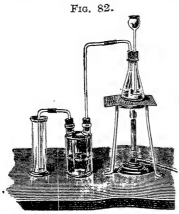
1396. Saturated Solution of Hydrogen sulphide (No. 31, 1410) is made by passing the gas from the apparatus described in par. 1371 or 1372 through water (Fig. 81) until the liquid is saturated (1395). The solution should be prepared in small quantity only, since the gas gradually undergoes decomposition, attended with deposition of sulphur, by the oxygen dissolved from the aim.

1397. Sulphurous acid Solution (No. 8, 1410) is prep-are d by passing SO₂ into cold water (1395). The gas may be made by heating copper turnings or clippings with strong H₂SO₄ in the flask shown in the apparatus in Fig. 82, and passing the gas through water in the washing-bottle.

This solution is kept in stock for pharmaceutical purposes, and may therefore be readily purchased. Cylinders of liquid 80, may also be purchased, and the gas which it evolves by spontaneus evaporation, when the valve is opened, may be employed for preparing the saturated solution.

1398. Chlorine—water (No. 22, 1410) is made by passing chlorine into cold water until the liquid is pale yellow in colour.

The apparatus for preparing the gas is shown in Fig. 82. Managanese dioxide in lumps is placed in the flask, and is covered with commercial hydrochloric acid diluted with an equal volume of water. The contents of the flask are then gently heated, and the



PREPARATION OF SULPHUR DIOXIDE OR CHLORINE.

gas is washed by passing it through a little water in the two-necked bottle.

The gas may also be obtained from a cylinder of liquid chlorime.

Chlorine - water is rapicly changed in daylight into so lution of hydrochloric acid, with escape of oxygen. The liquid must therefore be kept in a perfectly dark place, or in a bottle which is shielded from light by an opaque covering.

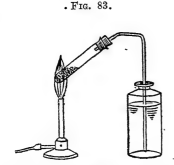
For many purposes brominewater (No.17, 1410) maybe substituted for chlorine-water, and is much more easily prepared and

preserved. Another substitute for chlorine-water is a little bleaching-powder shaken up with water in a test-tube and acidi fied with dilute HCl or HA.

1399. Nitrogen tetroxide Solution (No. 42, 1410)—Lead nitrate in dry powder is heated in a test-tube fitted with a cork and delivery-tube (Fig. 83). The reddish-brown gas which escapes is passed into dilute H₂SO₄.

Care must be taken to maintain the heat uniformly after the

fumes begin to be evolved, else the liquid may be sucked back into the hot tube by the contraction of the gas on cooling. For



SOLUTION OF NITROGEN TETROXIDE.



Passing Silicon Fluoride into Water.

the same reason the delivery-tube must be removed from the liquid as soon as the heating is stopped.

This solution need not be saturated with the gas.

• 1400. Hydrofluosilicic acid (No. 6, 1410) is made by passing silicon fluoride gas into water. The gas is prepared in a flask fitted as is shown in Fig. 83. An intimate mixture of 50 grams of dry, pure, finely-powdered fluorspar with 50 grams of fine white sand is poured into this flask; 300 grams of strong $\rm H_2SO_4$ are then introduced through the funnel tube, and the acid is thoroughly mixed with the powder by shaking it round in the flask.

SiF₄ is evolved when the flask is heated by a small flame from a rose-burner, or more safely on a sand-bath. The gas passes first through the two-necked bottle (Fig. 82), which must be empty and dry inside, thence through a bent tube ab (Fig. 84), which is fitted upon the outlet tube of the bottle by a rubber joint. The end of the tube (ab) dips into mercury contained in a small beaker d (Fig. 84), which stands in a large beaker containing 400 c.c. of water.

As soon as the silicon fluoride escapes from the mercury, and comes into contact with the water, it forms hydrofluosilicic acid which dissolves in the water, and silicic acid which remains suspended in the water as a gelatinous mass. The silicic acid would soon stop the escape of the gas, if the end of the delivery-tube were not protected from contact with water by being immersed in mercury.

When the current of gas slackens, the heat is raised until white fumes of H₂SO₄ begin to appear in the preparation flask. The process is then arrested, and the gelatinous matter is separated

from the liquid in the beaker. This is effected by squeezing the liquid through fine muslin, and then filtering, if necessary, to make it quite clear.

The silicic acid may be dried, heated strongly in a porcelain dish, and put by in a stoppered bottle for use as a reagent (No. 53, 1410).

STORE-JARS AND BOTTLES.

1401. Stock of Solid Chemicals.—The broad-necked common green-glass jars with flat stoppers, which are used for preserves and confectionery, may be used as store-jars for large quantities of solid chemicals. The jars are cheap and strong, and present the advantage over stoneware-jars that the quantity of material which they contain is seen at a glance. The flanged stoppers exclude dust and can be made air-tight by fastening a cork ring around their lower part. The complete exclusion of air is, however, seldom necessary.

These jars can be purchased in several convenient sizes. Solids are readily removed from them by means of wooden spoons or horn scoops.

- 1402. Solids for Analysis.—The 1 lb. glass honey-jars with glass caps serve well for keeping smaller quantities of solids, and can be made air-tight by means of a cork strip fastened round the neck. These jars serve for storing the solid powders which are required for analysis in the laboratory (1415–1419): their necks should be of sufficient size to allow of the insertion of an ordinary watch-glass to remove some of the contents.
- 1403. Reagents and Test-solutions.—Well-made white-glass bottles should be used for the Reagents and Test-solutions. The bottles should have flat-headed stoppers which protect the lip from dust, and each stopper should be well ground into the bottle so as to ensure its fitting. Stopper and bottle are prevented from permanently parting company by having an identical number etched upon them.

Accurately fitting glass stoppers should not be used in bottles containing solution of KOH- or NaOH-solution, since they are certain to be set fast. The use of a loosely fitting glass stopper, made to fit by surrounding it with a piece of rubber tube, obviates this difficulty.

The liquid bench reagents (1409) may be kept in narrow-necked bottles of 6 or 8 ounces (200 or 250 c.c.) capacity, and the solids in 4-ounce (120 c.c.) broad-necked bottles. The general reagents (1410) should be in 10 or 12-ounce (300 or 400 c.c.) bottles. The test solutions (1411, 1412) require larger bottles of about 32-ounce (1000 c.c.) capacity.

1404. Stock Bench-reagent Solutions.—The bench-reagent solutions (1409) may be conveniently kept in stock in an accessible position in the laboratory in large glass jars with glass taps below.

These taps may be ground into a tubulure near the bottom of the bottle, and after the perfectly dry tap has been greased and inserted, it is fastened in the tubulure by applying externally a thick coating of paraffin wax. The plug and seat of the stopper require to be occasionally perfectly dried and then greased. Or the taps may be inserted through a rubber cork which has been bored and placed in the tubulure. The storage of the bench-reagents in large stock-jars enables each student to fill his own bench-bottles when necessary.

1405. Stock-bottles for General Reagent- and Test-solutions.—These solutions may be made of double strength and may be kept in Winchester-quart bottles. When required for use a solution is diluted with an equal volume of water.

LABELLING BOTTLES.

1406. Every Bottle in use in the Laboratory should be Labelled.—Beginners are much assisted if the label bears not only the name but also the chemical formula of the substance which is contained in the bottle. It is also convenient to legibly number each bottle in the series. This number serves as a short means of reference. If the edges of the shelves are numbered to correspond with the bottles which should stand on them, the bottles are easily kept in their places and are quickly found when they are required.

The label should also give the strength of the solution in terms

of normality (1385).

The gummed labels should be stored in a dry place. When

the label is used, the whole of the gummed surface should be wetted and made to adhere to the bottle by pressure with a clean cloth. The label is then allowed to dry thoroughly, and is protected by being brushed rapidly across with an ordinary broad flat and stiff gum-brush, which has just been taken out of a dish of melted paraffin wax. A thin coating of the wax is thus made to entirely cover the label and to project for a short distance beyond its edge upon the glass surface.

A little practice will render it possible to lay on an even layer of the wax, which does not penetrate and stain the paper. If the paraffin is heated only slightly above its melting-point, it will set

too rapidly to penetrate the paper.

This paraffin varnish, if properly applied, perfectly resists the action of water, acids and alkalis. In fact, it can only be removed or injured by mechanical means, by heat, or by such solvents as ether and turpentine.

Bottles containing the strong mineral acids commonly have the name or formula etched upon their surface. This is costly and it is unnecessary, since the paraffined paper label resists the action even of strong acids.

Names and formulæ painted on bottles with Japan-black are also permanent in the laboratory.

LISTS OF REAGENTS AND CHEMICALS.

1407. The following paragraphs (1408-1414) give particulars of the reagents and chemicals required for the foregoing course of analytical work.

The chemicals required for Section II (1408), with a few exceptions, will be found by the reference figures in the later lists.

The chemical solutions and solids required for Section III fall into the following three classes:—

- 1. The Bench Reagents (1409) are those most frequently required for use by the student, and a set is therefore supplied to each working-bench. These bottles should be refilled when necessary by the student from the store-bottles (1404).
- 2. The General Reagents (1410) are not required so frequently; and one set, conveniently placed, will suffice for about fifteen students.
- 3. The Test-solutions and Solids (1411, 1412) are supplied for trying the reactions for metals and acid-radicles by means of the above reagents. There should be a set of these for every fifteen students in a convenient position in the laboratory. An (s) placed against the number means that the substance is required in the solid state as well as in solution: the solids may be in powder contained in 4-ounce (120 c.c.) wide-mouthed bottles.

It will be seen that the Bench Reagents, the General Reagents, and the Test Chemicals are placed in alphabetical order after the acids, each having an initial letter and a distinctive number.

The label of each bottle should carry this distinctive letter and number, and these should also be placed on the front edge of the shelf at the spot where the bottle should stand, as this keeps each bottle in its place and facilitates finding it. It is well also to state on the label of solutions the strength of the solution in the bottle.

The tabulated lists are arranged as follows:-

Column 1 gives the reference number and letter of the bottle.

Column 2 gives the name of the substance.

Column 3 gives the formula of the substance.

Column 4 gives the relation of the strength of the solution to normality (1385).

Column 5 gives the weight or volume of the substance to be contained in one litre of water.

Column 6 gives special information, when it is necessary.

The directions for making the solutions will be found in paragraphs 1384 et seq.

CHEMICALS REQUIRED FOR SECTION II.

1408. The following substances are required for the experiments in Section II.: they are placed in the order in which they are required for use by the student who is working through that section. When a reference number is affixed, it indicates that the substance will be found in one of the following lists. A letter (s) signifies that a soluble substance is required in the solid state.

It will be seen that only filter-paper, mercuric oxide, barium sulphate, wood charcoal and zinc oxide need be specially supplied, as the remaining substances are provided for in the subsequent lists.

Potassium nitrate (s)	
Copper sulphate (s) 12, 1411 Marble 39, 1412 Hydrochloric acid 2, 1409 Copper 24, 1410 Nitric acid 3, 1409	
Marble 39, 1412 Hydrochloric acid 2, 1409 Copper 24, 1410 Nitric acid 3, 1409	
Hydrochloric acid 2, 1409 Copper 24, 1410 Nitric acid 3, 1409	
Copper	
7, -4-7	
Barium chloride	
Ammonium carbonate 8, 1409	
Filter-paper	
Sulphuric acid 1, 1409	
Mercuric oxide	
Ammonium chloride (s) 2, 1411	
Potassium chloride (s)	
Sodium carbonate (s) 21, 1409	
Barium sulphate	
Manganese dioxide 38, 1410)
Borax (s)	
Ferrous sulphate (s) 14, 1411	
Sodium chloride	
Wood charcoal	
Lead acetate (s)	
Ammonium hydroxide 6, 1409	
Litmus-naner)	
Turmeric-paper 22, 1409)
Zinc oxide	
•	

1409. REAGENTS REQUIRED

The liquid reagents may be in 6 oz. (200 c.c.

small numbers								1	
	colur	Name. [The small numbers in brackets refer to remarks in column 6.]							Chemical Formula.
									*
phurie (¹)						•			H ₂ SO ₄
drochloric (2)					•••	•••			HCl
ric (3) .									HNO3
etic (4)				•••	٠.	•••	•••		HC2H3O2 or HA
ım chloride (5)									NH ₄ Cl
, or Ammonium	n hyd	iroxi	de (⁶)	•••	•••			NH40H
ım sulphide (7)									(NH ₄) ₂ S
ım carbonate (8	3) .				•••		•••		(NH ₄) ₂ CO ₃
ım oxalate .									(NH ₄) ₂ C ₂ O ₄ . H ₂ O
hloride				•••					BaCl ₂ .2H ₂ O
sulphate .									CaSO4
r Potassium hy	drox	ide (9)	•••					кон
n chromate									K2CrO4
n ferrocyanide				•••					K4FeCy6.3H2O
a ferricyanide (10)								K ₃ FeCy ₆
hosphate (11),	or .			•••		•••			Na ₂ HPO ₄ .12H ₂ O
mmonium phos	sphat	e							NaNH4HPO4.4H2O
arbonate							•••		Na ₂ CO ₃ .10H ₂ O (crystals)
)									Na ₂ B ₄ O ₇
n chlorate .					•••				KClO
n mixture (60)									Na ₂ CO ₃ +KCN
arbonate (13)									NagCO3
ers (51) .									
	arbonate (13).	arbonate (13)	arbonate (13)	arbonate (13)	arbonate (13)	arbonate (18)	arbonate (12)	arbonate (13)	arbonate (13)

FOR EACH BENCH.

4 oz (120 c.c.) bottles. bo

1			
	4.	5. Condition; and	6.
Reference	Strength	weight of solid in	Remarks.
Number.	in nor-	grams, or volume of liquid in cc. dis- solved to one litre.	[The small numbers refer to corresponding numbers in column 2.]
Liquids.			
1	3 N	83 c.c.	 Pure strong H₂SO₄ (Sp. gr. 1·84) is used; it must be gradually poured in the proper proportion into water contained
2	3 N	303 c.c.	in a large thin beaker or an earthen jar, while constantly
3	3 N	188 c.c.	stirring the water. The hot liquid is cooled by immersing the vessel in cold water, and is then poured into the store
4	3 N	840 c.c.	bottle. If the acid is pure, no white precipitate of PbSO ₄ will form during dilution.
5	3 N	160.5 grams.	 Pure strong HCl (Sp. gr. 1.16) is used; it must be colour- less, and give, after dilution, no precipitate with either BaCl₂
6	3 N	143 c.c.	or H2S, and no colour with KCNS-solution.
7	-	3500 c.c.	3. Pure strong HNO ₃ (Sp. gr. 1-42) is used; it must give no precipitate, after dilution, with BaCl ₂ - or AgNO ₃ -solu-
8	3 N	120 grams.	tion, added to separate portions, and no colour with KCNS- solution.
9	N	77 grams.	4. Pure acetic acid (Sp. gr. 1.04) is used; it must give no
10	N	122 grams.	precipitate or coloration with Am ₂ S. 5. Must give no precipitate with BaCl ₀ -solution.
11	N/20	Saturated (1393).	6. Must give only a very slight precipitate with lime-water,
12	3 N	120 grams.	no coloration with Am ₂ S, and separate portions acidified with HNO ₃ must give no precipitates with BaCl ₂ and
13	N	97 grams.	with AgNO ₃ -solution. "Liquor ammoniæ fortissima," of 880° specific gravity, is used for dilution with water.
14	N	606 grams.	 Must be yellow, and give with acids H₂S and a white pre- cipitate of S; it must give no precipitate with solutions
15		In small pieces.	of Ca- or Mg-salts.
	N	119 grams.	8. Solid Am ₂ CO ₂ is dissolved by the second method (1392) in cold water, but in diluting, one-fourth of the bottle
16	N	70 grams.	must be filled with strong AmOH. 9. White "stick potash" is used.
17	3 N	429 grams.	10. Dissolve a small piece of solid just before use; it must
Solids.			give no blue precipitate with FeCl ₃ -solution. 11. Its solution must yield no precipitate with AmOH.
18		In powder.	12. The borax is best dried by heating it in a platinum or
19		In small crystals.	gond: this solid mass is then milely powdered in a morear
20	_	In powder.	and kept in a stoppered bottle.
21	_	In powder.	13. Dry finely-powdered sodium carbonate. It must, after having been dissolved in excess of HNO ₃ , give no pre-
22	_	Blue and red lit mus, and turmeric, in small strips.	cipitate with BaCl ₂ , or AgNO ₃ , or AmHMoO ₃ -solution (571); and if evaporated with excess of HCl must leave no residue insoluble in dilute HCl (646).

1410. GENERAL REAGENTS, FOI These reagents may be contained in 10

1.			;	2.							3.
Reference	Name.							Chemical			
Number.	[The small numb	ers i	n bi	racke mn 6	ts re	fer t	o rei	nark	s in		Formula.
A 1	Acid, Hydrochloric (1)									HCl (Sp. gr. 1-16) .
A 2	Acid, Nitric (2)	•••	•••	***		•••	• • •	•••	• • •		
A 3	Acid, Sulphuric (3)										$H_2SO_4(Sp. gr. 1.84)$.
A 4	Acid Acetic (4)										
A 5	Acid, Hydrofluoric (5)									HF
A 6	Acid, Hydrofluosilici	c (6)								•••	H ₂ SiF ₆
A 7	Acid, Oxalic .										$H_2C_2O_4.2H_2O$
A 8	Acid, Sulphurous	•••	•••	• • •	•••	•••	•••	•••	•••	•••	H ₂ SO ₃
A 9	Acid Tartaric (7)										H2C4H4O6, or H2T .
A 10	Acid, Sodium tartra	te (8)	•••	•••	•••	•••	•••	•••	•••		$NaHC_4H_4O_6.H_2O$
A 11	Alcohol, Rectified sp	irit (9)								C2H6O
A 12	Ammonium acetate	•••	•••								NH ₄ C ₂ H ₃ O ₂
A 13	Ammonium chloride	(131	3)								NH ₄ Cl
A 14	Ammonium molybda										(NH ₄) ₂ MoO ₄
B 15	Barium carbonate (1	1)									BaCO:
B 16	Barium peroxide		•••								BaO ₂
B 17	Barium peroxide Bromine-water (12)				٠.					.	Br-water
C 18	Calcium carbonate (314)							,		CaCO
C 19	Calcium chloride										CaCl ₂ .6H ₂ O (crystals)
C 20	Calcium fluoride										AL 1985
C 21	Carbon disulphide										CS ₂
C 22	Chlorine-water										Cl-water
C 23	Cobalt nitrate (13)									.	Co(NO ₃) ₂ .6H ₂ O
C 24	Copper								•••		Cu
C 25	Copper sulphate									.	CuSO _{4.5} H ₂ O
E 26	Ether, methylated										
F 27	Ferric chloride (14)									.	FeCl _a .6H ₂ O
F 28	Ferrous sulphate (15)										
F 29	Fusion mixture (16)		•••	• • • •	***	•••	•••	•••	•••		$Na_2CO_3+K_2CO_3$.
G 30	Gold chloride	•••	•••	•	•						AuCl ₈
H 31	Hydrogen sulphide (17)									H ₂ S-water
I 32	Indigo-solution		•	•	•	•	•	•	•	,	

METALS AND ACID-RADICLES.

r 12 oz. (300 or 400 c.c.) bottles.

1	4.	5.	6.
	Strength	Condition; and	Remarks.
Reference	in nor-	weight in grams	[The numbers refer to corresponding small numbers in
Number.	mality (N)	dissolved to one	brackets in column 2.1
	(1385).	litre.	DATES OF THE ENGLISH BILL
	76.01	Strong, pure.	1. See remark 2 (1409).
A 1	10 N	Strong, pure.	2. See remark 3 (1409).
A 2	16 N		3. Must be colourless, and give no brown ring when it is
A 3	36 N	Strong, pure.	poured below some FeSO ₄ -solution (509).
A 4	3 N	Strong, pure	4. See remark 4 (1409).
A 5		_	5. Purchased and kept in gutta-percha bottles.
A 6		See par. 1400.	6. Must give no precipitate with Sr (NO2)2-solution.
A 7	. N	63 grams.	7. The crystallised acid must be freshly dissolved.
A 8	N/3	Saturated solu-	8. NaHT must be shaken with water in the bottle, in suffi-
		tion (1397). In small pieces.	cient quantity to leave a layer of solid below.
A 9	_	Saturated solu-	9. Strong rectified methylated spirit, which must leave no
A 10	_	tion.	residue on evaporation and must remain clear when it is
A 11		Strong.	diluted with water.
A 12	- N	77 grams.	10. 10 grams. of Am ₂ MoO ₄ are dissolved in 125 c.c. of strong AmOH, which has been previously mixed with an equa
A 13		In powder.	volume of water; the solution is then allowed to stand
A 14			if necessary until it is clear, and is then poured off into 500
B 15	_		c.c. of strong HNO3; the liquid will become hot, and mus-
B 16		In powder.	be cool before it is used.
B 17		_	11. Pure BaCO ₃ -powder is purchased, or is made by precipi
C 18		In powder.	tating BaCl ₂ -solution completely with Am ₂ CO ₃ -solution and washing by decantation. This powder is then mixed
C 19	N	110 grams.	with water to a creamy liquid.
C 20	1	In powder.	12. A few drops of Br are dissolved by shaking them up with
C 21		In powder.	water in the bottle. Sodium hypochlorite (Na ₂ Cl ₂ O), com
C 22	_	See par. 1398.	mercial "Liquor sodae chlorinatae" may be generally use
	N.	145 grams.	(311), but it does not keep well.
C 23	IN IN	Small strips of	13. The bottle should be closed by a loosely-fitting rubbe
C 24		sheet copper.	stopper, which is perforated to hold a glass tube passin
C 25	N	125 grams.	through and dipping into the liquid. By closing the ope- end of the tube with the finger and removing the tub
	14	120 81 112	some liquid is taken out, and a drop may be delivered b
E 26		90 grams.	carefully relaxing the pressure of the finger.
F 27	N.	In small pieces	1 11 11 11
F 28	-	TH SHIRL PIECES	stant stirring until the last drop produces a precipitat
F 29	-		of Fe(OH), and then filtering.
G 30	-	In dilute solu-	15. Clean small crystals; a small piece is crushed and dissolved in cold water immediately before use.
H 31	-	See par. 1396.	16. The dry finely-powdered carbonates are intimately
I 32	-	Indigo-carmine	mixed in the proportion of 53:69, and the mixture is kep
1 04		dissolved in	in a stoppered bottle.
	}	water.	17. The liquid must not darken when it is mixed with AmOH

1410. (Continued.) GENERAL REAGENTS FO

1.	2.	3.
Reference Number.	Name. [The small numbers in brackets refer to remarks in column 6.]	Chemical Formula.
L 33	Lead acetate	$Pb(C_2H_3O_2)_2.3H_2O$.
L 34	Lead acetate in potash (18)	PbA2+KOH
L 35	Lead nitrate	Pb(NO ₃) ₂
L 36	Lead nitrate Lime-water	1
M 37	Magnesium sulphate	MgSO ₄ .7H ₂ O
M 38	Manganese dioxide (19)	MnO _o
M 39	Marble	CaCO ₃
M 40	Manganese dioxide (10) Marble Mercuric chloride	
M 41	Microcosmic salt (20)	NaNH4HPO4.4H2O .
N 42	Nitrogen tetroxide solution	N2O4-solution
P 43	Microcosmic salt (*0) Nitrogen tetroxide solution Platinum chloride (Note, 420)	PtCl4
P 44	Potassium chloride	KCI
P 44 P 45	Potassium cyanide (21)	KCN
P 46 47 48 49 PP 551 552 554 555 555	Potassium hydrogen sulphate Potassium ierricyanide Potassium nitrate Potassium nitrate Potassium nitrite (22) Potassium thiocyanate Silica, white sand (1400) Silver nitrate	
S 56 S 57	Sodium acetate	Na Land Wi
S 58	Sodium hydroxide, pure (25)	NaOH
S 59	Sodium acetate and acetle acid (24) Sodium hydroxide, pure (25) Sodium nitrate, solid	NaNO ₃
S 60 S 61 S 62 W 63 Z 64	•	SnCl ₂ .2H ₂ O Fe
Z 65		ZnSO4.7H2O

METALS AND ACID-RADICLES.

Reference Number	Strength in nor- mality (N) (1385).	5. Condition; and weight in grams dissolved to one litre.	6. Remarks. [The numbers refer to corresponding small numbers in brackets in column 2.]
L 33	N	190. Clear by adding HA.	18. Add to some of the above warm PbA2-solution (L 33)
L 34 L 35	_	Small pieces.	KOH-solution (12, 1409) with constant stirring, until the precipitate just redissolves. Prepare in small quantity
L 36	N/20	Saturated solu- tion (1393).	19. Fine powder; must not evolve CO ₂ or Cl when it is
M 37	N	123 grams. Solid, in powder	warmed with strong H ₂ SO ₄ . 20. The solution in HNO ₃ must give no precipitate with
M 38 M 39		Pea-sized pieces.	AgnO ₃ -solution.
M 40 M 41	N/5	27 grams. In powder.	 Dissolve a small piece by warming it with water immediately before use.
N 42 P 43	. =	See par. 1399. [Costly, use with	22. Make in small quantity only, or immediately before use, as the solution changes. It must give abundant
P 44	N	care.] 75 grams.	red fumes when mixed with dilute H ₂ SO ₄ .
P 45		In small pieces or powder.	23. Freshly burnt pieces of lime are placed upon a plate, and water is poured upon them until they begin to appear
P 46 P 47	N	49grms. Andsolid Small pieces.	moist on the surface; the superfluous water is then drained off, and as soon as the lime has crumbled to powder and
P 48		Small pieces.	cooled, it is placed in a broad-mouthed stoppered bottle. 24. Dissolve 20 grams of sodium acetate crystals in 60 c.c. of
P 49 P 50	. N/10	17 grams. In small pieces.	distilled water, and add 40 c.c. of strong HÅ (A 4).
P 51 P 52	N N	85 grams. 97 grams.	25. This solution should be kept in a green glass bottle as it dissolves Pb from colourless flint glass; and a rubber or
S 53 S 54	N/10	In fine powder. 17 grams.	loose stopper is preferable to an ordinary ground-in glass stopper, as this becomes fixed. The solution must not
S 55 S 56	N	In powder. 65 grams	darken in colour with H ₂ S, or give gelatinous Al(OH) ₃ when it is mixed with AmCl-solution (1409, 5).
S 57 S 58	3 N	120 grams.	26. SnCl ₂ -crystals are dissolved in water containing HCl, and the solution is kept with immersed tin fragments and
S 59		In small crystals,	excluded from air.
S 60	N	112.5 grams.	 Make before use by stirring 2 grams of starch-powder, made into paste with water, into 100 c.c. of boiling water,
S 61 S 62		In powder.	and cool.
	_	In pieces.	28. Short lengths of broken knitting-needles, kept with quick-lime to prevent rusting.
W 63 Z 61		Strips, rods, or granulated.	29. Must be free from arsenic (176, 177, 178).
Z 65	N	145 grams.	20. Many 20 1100 Minetito (1/0, 1//, 1/0).

1411. SOLUTIONS FOR THE

These solutions may be kept in 32 oz. (1000 c.c.) bottles, each of which is labelled Each substance with (s) affixed in column 1 should also be provided as a solid in

1.			2								3.
Reference		Name.								Chemical Formula.	
Number.										- 1	Formula.
	[Small numbers	in b c	racke	ets r m 6.	eter]	to re	marı	is in			
A [1.s	Aluminium potassium	su	lphat	e.							AlK(SO ₄) ₂ .12H ₂ O
A 2.s	Ammonium chloride				***	***	***	***	***	}	Amcl
A 3.s	Antimonious chloride	(1)					. "		•		SbCl ₃
A 4.s	Arsenious oxide in H	C1 (†	²)				***	***	***		As406
A 5	Arsenious oxide in H	0 (3)						•		As406
B 6	Barium chloride				•••	***	• • •	***	•••	***	BaCl ₂ .2H ₂ O
B 7.s	Bismuth nitrate (4)									- 1	Bi(NO ₃) ₃ .5H ₂ O
C 8.8	Cadmium sulphate Calcium chloride	***		***		• • •		***	• • •	***	CdSO4.4H2O
C 9	Calcium chloride .							•:			CaCl ₂ .6H ₂ O
C 10.s	Chromium potassium	sul	pnate	3		• • •	***	***	• • • •	***	CrK(SO ₄) ₂ .12H ₂ O
C 11	Cabalt mitrata								•		Co(NO ₃) ₂ .6H ₂ O
C 12.5	Copper sulphate		***			***			•••	***	CuSO _{4.5} H ₂ O
F 13	Ferric chloride .								-		FeCl ₃ .6H ₂ O
F 14.5	Ferrous sulphate (5)				•••				***	• • •	FeSO4.7H2O
L 15.s	Lead acetate (°) .					•	4		•	•	Pb (C ₂ H ₃ O ₂) ₂ .3H ₂ O
M 16.s	Magnesium sulphate						•••		***	• • •	MgSO ₄ .7H ₂ O
M 17.s	Manganese chloride								^ •		MnCl ₂
M 18.s	Mercuric chloride		•••			• • •		• • •		***	HgCl2
M 19	Mercurous nitrate (7)										$Hg_2(NO_3)_2.H_2O$
N 20.s	Nickel sulphate							• • •	***		NiSO4.7H2O
P 21.s	Potassium chloride										KCl
S 22	Silver nitrate							***		• • •	AgNO ₈
S 23.s	Sodium arsenate .										Na ₂ HAsO ₄ .12H ₂ O
S 24.8	Sodium chloride					•••		***	***	• • •	NaCl
S 25	Stannic chloride (8)										SnCl ₂ .2H ₂ O
S 26.s	Stannous chloride (*)		***					***	***	***	SnCl ₂ .2H ₂ O
S 27	Strontium nitrate										$Sr(NO_3)_2.4H_2O$
Z 28.s	Zinc sulphate				***		***		•••		ZnSO _{4.7} H ₂ O

REACTIONS OF THE METALS.

1411]

with the number, name, formula and normality-strength of the solution (1385), a 4 oz. (120 c.c.) wide-mouthed bottle similarly labelled.

	4.	5	6.
Reference Number	Strength in nor-	Weight in grams	Remarks.
Mumbu	mality (N). (1385).	dissolved in one litre of water.	[The numbers refer to corresponding small numbers in brackets in column 2.]
A 1.s	N	118.	1. 5 grams of SbCl ₃ crystals are dissolved in 25 c.c. of strong
A 2.s A 3.s	N N/5	54. 15.	HCl diluted with 25 c.c. of water, and the solution is diluted to a litre with strong HCl diluted with four volumes of
A 4.3 A 5	N/5	7. Saturated.	water. For tests on the solid, Sb ₂ O ₃ or SbOCl is used.
В 6	O N	122.	2. Dissolve the As ₄ O ₆ by heating it with 25 c.c. of strong HCl diluted with 25 c.c. of water, and dilute to a litre.
B 7.s C 8.s	$\frac{N/5}{N/5}$	30. 26.	3. Boil the As ₄ O ₆ for some minutes with 1000 c.c. of water
C 9	N/2	55.	and filter into the bottle.
C 10.8 C 11	N/5 N/5	20. 29.	4. The Bi(NO ₃) ₃ is dissolved by heating it with 25 c.c. of strong HCl diluted with 25 c.c. of water: this is cooled
C 12.s	N/5	25.	and poured into the bottle, and the volume is made up
F 13	N/2 N/5	45. In small pieces.	with water containing 20th its volume of strong HCl. For the solid, BiONO2 is used.
F 14.s L 15.s	N/5	38.	5. A small crystal of FeSO ₄ (F 28, 1410) is crushed and dis-
M 16.s	N	13. 40.	solved in cold water just before it is used. 6. Some HÅ must be added to clear the solution.
M 17.s M 18.s	N/5 N/5	29.	7. Dissolve 56 grams of Hg ₂ (NO ₃) ₂ by warming it with 6
M 19	N/5	56.	c.c. of strong HNO ₃ diluted with 114 c.c. of water, then
N 20.s P 21.s	N/5 N	28. 75.	dilute with water to a litre. Keep drops of Hg at the?
S 22	N/5	34.	8. Heat 28 grams of SnCl ₂ with 25 c.c. of strong HCl and 100
S 23.s S 24.s	N/5 N	27. 58.	c.c. of water, adding powdered KClO ₃ gradually with constant stirring, until the liquid turns yellow and Cl is smelt:
S 25	N/2	45.	then boil off the Cl, and dilute to one litre.
S 26's S 27	N/5 N	23. 141.	9. Boil 23 grams of SnCl ₂ with 25 c.c. of strong HCl diluted: with 25 c.c. of water, and dilute the clear solution to one?
Z 28.s	N/5	29.	litre. Keep scraps of tin in the bottle.
	ŀ	1	

1412. SOLUTIONS FOR THE

These solutions may be kept in 32 oz. (1000°c.c.) bottles, each of which is labelled Each substance with an (s) affixed in column 1 should also be provided as a solid ist occur in the preceding tables, as is shown by the references in square brackets reference; but it is more convenient to make up this set complete, as below, if themical name is placed before the metal in this list, the ordinary chemical name

A 1.s Acetate, sodium [1. Reference Number.	2. Name. [Small numbers in brackets refer to column 6. References in square brackets to preceding Tables.]	3. Chemical Formula.
T 28.s Tartaric acid	8.8.8.8.8.8.8.9.01123445678901123445678901123456789012322222222222222222222222222222222222	Arsenate, sodium Borate, sodium Borate, sodium Borate, sodium Bromide, sodium Carbonate, calcium (marble) Carbonate, calcium (marble) Carbonate, sodium Carbonate, sodium Clorate, potassium Chloride, sodium (*) Chromate, potassium Cyanide, potassium Cyanide, potassium Cyanide, potassium Ferricyanide, potassium Fluoride, calcium (fluorspar) Fluosilicie acid Hypochlorite, sodium (*) Indide, potassium Nitrate, potassium Nitrate, potassium Nitrate, potassium Nitrate, potassium Nitrate, sodium Silicate, ammonium Fluoride, sodium Silicate, sodium (*) Silicate, sodium (*) Silicate, sodium Sulphide, sodium Sulphide, ammonium Sulphide, ferrous Sulphite, sodium Tartrate acid Tartrate, sodium	Na.HASO.12H2O Na.2B4O.10H2O Na.2B4O.10H2O Na.2B4O.10H2O Na.2B4O.3 Na.Cl

REACTIONS OF THE ACID-RADICLES.

with the number, name, formula and normality-strength (1385) of the solution. in a 4 oz. (120 c.c.) wide-mouthed bottle similarly labelled. Many solutions in this in column 2: these may be used if necessary from the bottles indicated by the there is sufficient shelf-room in the laboratory. The acid-radicle portion of the being inverted, in order to arrange the acid-radicles alphabetically.

Reference Number. Condition, and strength in normality (N) (1385). Grams dissolved to one litre one litre numbers refer to corresponding small numbers in brackets, column 2.1	
11: 1: 1: 1	
A 1	3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3

1413. Special Test Substances and Reagents for the Rarer Metals and Acid-Radicles.

$\begin{array}{cccccccccccccccccccccccccccccccccccc$	H. ACID-RADICLES. Group III:— Bromate
Group II:— Palladium chloride Gold chloride [30, 1410] Platinum chloride [43, 1410] Platinum chloride NæglrClg	K Mn()
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
$ \begin{array}{llllllllllllllllllllllllllllllllllll$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
Thorium sulphate Th(SO ₄) ₂	III. REAGENTS.
Indium sulphate $UO_2(NO_3)$ Uranyl nitrate $UO_2(NO_3)$ Vanadate $NaVO_3$	Potassium bicarbonate solu- tion KHCO:
Group V:— Lithium chloride . LiCl Rubidium chloride . RbCl Caesium chloride . CsCl	Hydrogen peroxide, dilute acid solution H ₂ O ₂ Calcium sulphate, fine powder. CasO ₄

1414. TEST SUBSTANCES AND REAGENTS FOR ORGANIC SUBSTANCES.

Reagents already in use for inorganic substances are not included.

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ORGANIC SUBSTANCES.
                                                                                                                              Group VII. Glucosides :--
Group I. Cyanogen Acid-radicles :--
                                                                                                                                                                                    \substack{\substack{C_{13}H_{18}O_7\\C_{29}H_{46}O_{12}\\C_{20}H_{27}O_{11}N}
                                                                                                                                    Salicin
      Cyanide . . . KCN
                                                                                                                                    Digitalin
                                              . KČNO
      Cyanate .
                                                                                                                                    Amygdalin .
                                                     KCNS
     Thiocyanate
                                              K<sub>4</sub>FeCy<sub>6</sub>.3H<sub>2</sub>O
K<sub>3</sub>FeCy<sub>6</sub>
     Ferrocyanide.
     Ferricyanide
                      2. Aliphatic Acid-radicles :-
                                                                                                                               Group VIII. Organic bases :-
     2. Aliphatic Acid-radicles:—
Formate . NaCHO<sub>2</sub>
Acetate . NaC<sub>2</sub>H<sub>3</sub>O<sub>2.3</sub>H<sub>2</sub>O
Oxalate . (NH<sub>4</sub>)<sub>2</sub>O<sub>5</sub>O<sub>4.2</sub>H<sub>2</sub>O
Succinate . (NH<sub>4</sub>)<sub>2</sub>O<sub>5</sub>O<sub>4.2</sub>H<sub>2</sub>O
Malate . H<sub>2</sub>O<sub>4</sub>H<sub>4</sub>O<sub>6</sub>
Tartrate . KNaC<sub>4</sub>H<sub>4</sub>O<sub>6</sub>, and
NaHC<sub>4</sub>H<sub>4</sub>O<sub>6</sub>
Citrate . Na<sub>2</sub>O<sub>6</sub>H<sub>5</sub>O<sub>7.2</sub>H<sub>2</sub>O
                                                                                                                                    Aniline . .
                                                                                                                                                                                     {^{{
m C}_6{
m H}_5.{
m NH}_2}_{{
m C}_6{
m H}_4.{
m CH}_3.{
m NH}_2}}
                                                                                                                                     Toluidine
                                                                                                                                                                                    C<sub>6</sub>H<sub>4</sub>.CH<sub>5</sub>.N H<sub>2</sub>
C<sub>5</sub>H<sub>5</sub>N
C<sub>9</sub>H<sub>7</sub>N
C<sub>6</sub>H<sub>5</sub>.NH.CO.CH<sub>5</sub>
C<sub>8</sub>H<sub>5</sub>O.C<sub>6</sub>H<sub>4</sub>.NH.COCH<sub>7</sub>
C<sub>11</sub>H<sub>12</sub>N<sub>2</sub>O
CO(NH<sub>2</sub>)<sub>2</sub>
                                                                                                                                    Pyridine .
                                                                                                                                     Quinoline.
                                                                                                                                    Acetanilide .
Phenacetin .
                                                                                                                                     Antipyrin
                                                                                                                                     Urea . .
                      3. Aromatic Acid-radicles :-
                                                     : Acta-rataces: —

NH4C,H5O<sub>2</sub>

NAC,H5O<sub>3</sub>

HC,H<sub>5</sub>O<sub>3</sub>

HC,H<sub>5</sub>O<sub>3</sub>

HC,H<sub>5</sub>O<sub>3</sub>

HC<sub>2</sub>H<sub>2</sub>O<sub>7</sub>,3H<sub>2</sub>O

KC<sub>3</sub>H<sub>3</sub>N<sub>4</sub>O<sub>3</sub>
      Benzoate
     Salicylate.
Gallate
                                                                                                                               Group IX. Alkaloids :-
                                                                                                                                                                                   48:—

C20H24N2O2.3H2O

2C20H24N2O2.5H2O

C19H23N2O

C19H23N2O

C17H19NO3

C17H17NO.0CH3.OH

C22H22NO7

C20H21NO4

C21H22N2O2

C23H2CN2O4.4H2O

C3H00N4O2.H2O

C17H21NO4

C22H42NO5

C17H21NO4

C22H42NO7

C3H20N2O2.4H2O

C3H20N2O4.4H2O

C3H20N2O3.0H2O
      Tannate
                                                                                                                                      Quinine . .
      Hippurate
                                                                                                                                      Quinidine.
      Meconate . . .
                                                                                                                                      Cinchonine .
Urate . . KO<sub>5</sub>H<sub>3</sub>N<sub>4</sub>O<sub>3</sub>
Group II. Hydrocarbons and their Deriva-
                                                                                                                                     Morphine.
                                                                                                                                      Apomorphine
                                                tives :-
                                                                                                                                     Codeine . . .
                                                       C<sub>10</sub>H<sub>16</sub>
C<sub>6</sub>H<sub>6</sub>
C<sub>10</sub>H<sub>8</sub>
C<sub>14</sub>H<sub>10</sub>
      Turpentine .
      Benzene .
     Benzene . .
Naphthalene
Anthracene .
                                                                                                                                      Papaverine
                                                                                                                                      Strychnine .
                                                                                                                                     Chloroform
                                                       CHCl,
      Carbon tetra-
chloride
                                                                                                                                      Cocaine . . .
Veratrine . .
                                                       CHI,
                                                                                                                                                                                      C<sub>17</sub>H<sub>23</sub>NO<sub>3</sub>
C<sub>17</sub>H<sub>23</sub>NO<sub>3</sub>
C<sub>17</sub>H<sub>21</sub>NO<sub>4</sub>
      Iodoform .
                                                                                                                                      Atropine .
      Carbon disulphide CS2
                                                                                                                                     Hyoscyamine
Group III. Alcohols and Ethers:—
Methyl alcohol . CH<sub>2</sub>OH
Ethyl alcohol . C<sub>2</sub>H<sub>5</sub>OH
                                                                                                                                      Hyoscine
                                                                                                                                     Nicotine .
                                                                                                                                                                                     C<sub>10</sub>H<sub>14</sub>N<sub>2</sub>
C<sub>8</sub>H<sub>17</sub>N
                                                                                                                                      Coniine .
                                                       C<sub>5</sub>H<sub>1</sub>OH
C<sub>3</sub>H<sub>5</sub>(OH)<sub>3</sub>
(C<sub>2</sub>H<sub>5</sub>)<sub>2</sub>O
C<sub>2</sub>H<sub>5</sub>.C<sub>2</sub>H<sub>3</sub>O<sub>2</sub>
      Amyl alcohol
      Glycerol . .
Ethyl ether .
      Ethyl acetate
Group IV. Phenol and its Derivatives:—Phenol . . . CeHs.OH
                                                       C<sub>6</sub>H<sub>5</sub>.OH

C<sub>6</sub>H<sub>4</sub>(OH)<sub>2</sub>

C<sub>6</sub>H<sub>3</sub>(OH)<sub>3</sub>

C<sub>6</sub>H<sub>2</sub>(NO<sub>3</sub>)<sub>3</sub>OH
      REAGENTS.
      Pyrogallol . . Pieric acid . .
                                                                                                                                Sodium picrate, satu-
 Group V. Aldehydes and Ketones:-
                                                                                                                                     rated solution
                                                                                                                               neutral solution . Hg(NO<sub>2</sub>)<sub>2</sub>
Quicklime, in powder CaO
Potassium bicarbon-
ate solution
                                                                                                                                Mercuric nitrate,
neutral solution .
            (solution) .
                                                        H.COH
                                                       CH<sub>s</sub>.COH

CCl<sub>s</sub>.CH(OH)<sub>2</sub>

C<sub>6</sub>H<sub>s</sub>.COH

CO(CH<sub>s</sub>)<sub>2</sub>
       Acetaldehyde
       Chloral hydrate .
      Benzaldehyde .
                                                                                                                               Iodic acid, solution
(1:15)
Iodine, in small
fragments
 Acetone . . . CO(CE
Group VI. Carbohydrates:-
                                                                                                                                                                                      HIO,
                                                        C_6H_{12}O_6
       Dextrose . . .
                                                      C<sub>6</sub>H<sub>12</sub>O<sub>6</sub>
C<sub>12</sub>H<sub>22</sub>O<sub>11</sub>
C<sub>12</sub>H<sub>22</sub>O<sub>11</sub>.H<sub>2</sub>O
C<sub>12</sub>H<sub>22</sub>O<sub>11</sub>.H<sub>2</sub>O
       Levulose .
                                                                                                                                 Yeast
       Sucrose .
                                                                                                                                Yeast . . . Rochelle salt
                                                                                                                                                                                      KNaT.4H.0
       Maltose .
                                                                                                                                 Rosaniline (Fuchsine)
      Lactose .
                                                                                                                                                                                      ZnCl_2
                                                                                                                               Zinc chloride
Ammonium iodide
Sodium nitroprusside
                                                       (C_6H_{10}O_5)n
(C_6H_{10}O_5)n
       Cellulose .
                                                                                                                                                                                      NH4I
       Starch
     Starch .
Dextrin .
                                                        (C6H10O5)n
```

SUBSTANCES TO BE GIVEN FOR ANALYSIS BY THE PRECEDING TABLES.

It must be understood that the substances mentioned below are merely brought forward as examples of what may be given to the student for analysis. The teacher will use his own judgment in adopting and extending the list.

SUBSTANCES FOR ANALYSIS WHILE TRYING THROUGH THE REACTIONS FOR METALS AND ACID-RADICLES.

1415. After working through the reactions for a Group of Metals or Acid-radicles, a few unknown substances containing only one member of the Group are tested; the number of members present is then gradually increased in the substances which are subsequently given.

It is well to keep these substances in the dissolved state (1411, 1412), since they are intended mainly to afford practice in separation and detection according to the Group-tables; the time spent by the student in preparing the solution would therefore be wasted.

Occasionally, however, a solid substance may be given, in which the Metal or Acid-radicle present is to be detected by blowpipe tests, or by other tests made on the solid substance. The following will serve as examples of solids:—

Group V. Solution of NaCl " NH ₄ Cl " KCl " NACl+KCl " NACl+NH ₄ Cl+ KCl " NACl+NH ₄ Cl+ KCl " MgSO ₄ .7H ₂ O+ NaCl+KCl Solids: NaCl, NH ₄ Cl, KCl, AmCl+NaCl, etc.	$\begin{array}{c} & & & & & & & & & \\ & & & & & & & & \\ Solution \ of \ BaCl_2.2H_2O \\ "" & Sr(NO_3)_2.4H_2O \\ "" & ScCl_2.6H_2O \\ "" & Sr(NO_3)_2.4H_2O \\ "" & Sr(NO_3)_2.4H_2O \\ "" & BaCl_2.2H_2O + \\ "" & Sr(NO_3)_2.4H_2O \\ "" & Sr(NO_3)_2.4H_2O \\ "" & Sr(NO_3)_2.4H_2O \\ \end{array}$	Group III.b. Solution of ZnSO ₄ .7H ₂ O "
---	--	---

SIMPLE SUBSTANCES FOR ANALYSIS.

1416. Any one of the solutions named in paragraphs 1411, 1412 may be given for analysis: some of these are neutral, some acid, and others alkaline in reaction. The following more difficult solutions, with acid or alkaline reaction, may also be added to the list:—

Acid.	Alkaline,
$Ca_3(PO_4)_2$, "Bone-ash" dissolved in dilute HCl. BaC_2O_4 dissolved in dilute HCl. $Mg(BO_2)_2$, " HCl. $BaCrO_4$, " HCl.	KSbO ₃ dissolved in water. Na ₂ SiO ₃ , solution of "soluble glass." Na ₂ SnO ₃ , " preparing salt" of the dyer.

*Made by mixing hot solutions of Na₂B₄O₇ and Mg(NO₃).

As examples of substances possessing metallic appearance (1299), the following may be mentioned as suitable for analysis:—

Zn, in pieces or filings.

Pb ,, ,, Graphite, or Black lead.

FeS₂, Iron pyrites.

Iron filings.

As insoluble substances which require to be examined by paragraphs 1306, et seq., the following may be given for analysis:—

BaSO₄.

PbSO₄.

AgCl.

SnO₂, Tinstone.

CaF₂, Fluor spar.

FeCr₂O₄, Chrome Iron ore.

S, as pieces of roll sulphur.

C, as powdered wood charcoal or plumbago.

• SOLID SUBSTANCES TO BE ANALYSED BY THE PRELIMINARY EXAMINATION IN SECTION VI.

1417. For the Preliminary Examination for Metals (1215-1224):—

	Simple.		Com	plex.
NH ₄ Cl HgCl ₂ As ₄ O ₆ KNO ₃ Cr ₂ O ₃	$\begin{array}{c} {\rm Pb(NO_3)_2} \\ {\rm NaCl} \\ {\rm BaCl_2.2H_2O} \\ {\rm Sr(NO_3)_2.4H_2O} \\ {\rm CuSO_4.5H_2O} \\ {\rm (See} \ {\it Note.}) \end{array}$	$\begin{array}{c} \rm ZnSO_4.7H_2O\\ MnCl_2\\ \rm Co(NO_3)_2.6H_2O\\ NiSO_4.7H_2O\\ SnO_2 \end{array}$	$\begin{array}{c} {\rm AmCl+NaCl} \\ {\rm HgCl_2+BaCl_2.2H_2O} \\ {\rm MnCl_2+KCl} \\ {\rm NiSO_4.7H_2O} + \\ {\rm ZnSO_4.7H_2O} \end{array}$	$\begin{array}{c} {\rm AlK(SO_4)_2.12H_2O} \\ {\rm SnO_2 + Sr(NO_3)_2.4H_2O} \\ {\rm CaCO_3 + ZnSO_4.7H_2O} \\ {\rm Cr_2O_3 + MnCl_2 +} \\ {\rm NaCl} \end{array}$

Note.—Best finely powdered, since its colour is then almost destroyed.

For the Preliminary Examination for Acid-radicles (1225-1228):—

	Complex.			
CaCO ₃ Na ₂ SO ₃ .7H ₂ O CaCl ₂ O NaCl	FeS KNO ₃ KClO ₃ KI	Na ₂ S ₂ O ₃ .5H ₂ O KBr Na Å	$\begin{array}{c} \operatorname{CaCO_3} + \operatorname{KNO_3} \\ \operatorname{CaCl_2O} + \operatorname{CaF_2} \\ \operatorname{Na_2CO_3} + \operatorname{KI} \end{array}$	

A few of the above-mentioned solids may then be examined by both Preliminary Examinations for Metals and for Acid-radicles.

Substances to be analysed by the General Course in Section VI.

1418. The following lists are so arranged that the analysis of the substances contained in them is more difficult in each column proceeding from left to right, and usually from top to bottom in one and the same column. For examples of alkaline solutions, see paragraph 1416.

1. Simple soluble substances.	2. Complex soluble substances.	3. Complex soluble substances.
BaCl ₂ .2H ₂ O MnCl ₂ HgCl ₂ MgSO ₄ .7H ₂ O KCl NiSO ₄ .7H ₂ O *CaCO ₃ Pb(NO ₃) ₂ *As ₄ O ₆ FeCl ₃ NH4Cl NaCl etc., etc.	$\begin{array}{c} {\rm CaCl_2.6H_2O + BaCl_2.2H_2O + Sr(NO_3)_2.4H_2O} \\ {\rm AlK(SO_4)_2.12H_2O + CrK(SO_4)_2.12H_2O} \\ {\rm MnCl_2 + ZuSO_4.7H_2O + CuSO_4.5H_2O} \\ {\rm Pb(NO_3)_2 + HgCl_2 + NaCl} \\ {\rm *As_4O_4 + CaCO_3 + BaCO_3} \\ {\rm *ZnO + MgCO_3 + BaCO_3} \\ {\rm *MgCO_3 + BaCO_3 + PbCO_3} \\ {\rm etc., etc.} \end{array}$	FeCl ₃ MnCl ₂ ZnSO ₄ .7H ₂ O KCl

^{*} To be given in the solid state.

•	Substances yielding a Phosphate-precipitate in Group III.	5. Substances partly or entirely insoluble.	Metallic substances (1299-1305) :—
	{ CaCl ₂ .6H ₂ O FeCl ₂ Na ₂ HPO ₄ .12H ₂ O BaCl ₂ .2H ₂ O MgCl ₂ FeCl ₂ Na ₂ HPO ₄ .12H ₂ O	$\begin{array}{c} {\rm BaSO_4} \\ {\rm SiO_2 + S} \\ {\rm BaSO_4 + CaF_2 + SnO_2} \\ \\ \left\{ \begin{array}{c} {\rm BaCl_2 2H_2O} \\ {\rm AIK(SO_4)_2.12H_2O} \\ {\rm AgNO_3} \end{array} \right. \end{array}$	Iron pyrites (FeS ₂) Iron filings (Fe) Zinc clippings (Zn) Brass filings (Cu+Zn) Nickel silver (Cu+Zn+Ni) Bronze (Cu+Sn) Type-metal (Sb+Sn+Pb) Aluminium bronze (Cu+Al)
	(Na ₂ HrO ₂ .12H ₂ O (CaCl ₂ .6H ₂ O (CaCl ₂ .6H ₂ O MnCl ₂ CTK(SO ₄) ₂ .12H ₂ O Na ₂ HPO ₄ .12H ₂ O etc., etc.	$\begin{cases} \overset{\text{SiO}_2}{\text{S}} \\ \overset{\text{S}}{\text{C}} \end{cases}$ $PbSO_4 + BaSO_4$ $PbCrO_4, \text{ ignited.}$	Silicates (1312-1315):— Kaolin (Al ₂ O ₂ .2SiO ₂) Brown-clay (do. +Fe) Window-glass (Na, Ca, SiO ₂) Flint-glass (Pb, K, SiO ₂) Cyanogen Compounds (1316-1322):
	•		—Any of the foregoing which has been mixed with KON, KCNS, K ₄ FeCy ₆ , 3H ₂ O, K ₂ FeCy ₆ , K ₂ CoOy ₆

MINERALS FOR ANALYSIS

1419. The minerals are ranged under one of the principal metals which they contain, and these metals are placed in the order in which they occur in the Analytical Reactions (101-448). The names of minerals which contain the less common metals will usually be found above the reactions of each of those metals. In this list only the more common natural compounds are included, and the traces of impurities which they may contain are not specified in stating their composition by formulae.

GROUP I.—SILVER GROUP.

Lead :-

Galena, PbS. Cerussite, PbCO₃. Anglesite, PbSO₄. Leadhillite, PbSO₄.3PbCO₃. Pyromorphite, 3Pb₃(PO₄)₂.PbCl₂. Crocoisite, PbCrO₄.

Silver :-

Native silver, Ag. Horn silver, AgCl. Silver glance, or Argentite, Ag₂S. Pyrargyrite, Ag₃SbS₃. Proustite, Ag₃AsS₃.

GROUP II.A.—COPPER GROUP

Mercury :---

Cinnabar, HgS.

Bismuth:

Native bismuth, Bi. Bismuth ochre, Bi₂O₃. Bismuthine, or Bismuth glance, Bi₂S_{3*} Complex bismuth sulphides.

Copper:-

Native copper, Cu.

Cuprite, or Red copper ore, Cu₂O.

Copper glance, Cu₂S.

Copper pyrites, or Peacock copper ore, CuFeS₂.

Ernbescite, or Bornite, Cu₃FeS₃.

Tetrahedrite, 4Cu S.Sb S.

Malachite, CuCO₃.Cu(OH)₂.

Azurite, or Chessylite, 2CuCO₃.Cu(OH)₂.

Cadmium :-

Greenockite, CdS.

GROUP II.B.—ARSENIC GROUP.

Arsenic :--

Native arsenic, As.

Realgar, As₂S₂.

Orpiment, As₂S₃.

Mispickel, FeAsS.

Copper-nickel, NiAs.

Arsenical nickel, NiAs2.

Arsenosulphides.

Antimony :-

Native antimony, Sb.

Stibnite, or Grey antimony, Sb₂S₃.

Complex antimony sulphides.

White antimony ore, Sb₂O₃.

Red antimony ore, Sb₂O₃.2Sb₂S₃.

Tin :-

Tinstone, or Cassiterite, SnO₂.

Stannite, or Tin pyrites, Cu₂FeSnS₄.

GROUP III.A.-IRON GROUP.

Aluminium :-

Corundum, Al₂O₃.

Bauxite, Al₂O₃(Fe₂O₃).2H₂O.

Diaspore, Al₂O₃.H₂O.

Alumstone, $K_2O.3Al_2O_3.4SO_3.6H_2O$.

Cryolite, AlF₃.3NaF.

Felspar (see K and Na).

Clay, Al₂Si₂O₇.2H₂O.

Iron:

Meteoric iron, Fe + Ni.

Red haematite, or Specular iron ore, Fe₂O₃.

Brown haematite, or Limonite, 2Fe₂O₃.3H₂O.

• Magnetite, or Magnetic iron ore, Fe₃O₄.

Iron pyrites, FeS2.

Pyrrhotite, or Magnetic pyrites, Fe₁₁S₁₂.

Chalybite, or Spathic iron ore, FeCO₃.

Clay ironstone, FeCO₃ + clay.

Black-band ironstone, $FeCO_3$ + bitumen.

Chromium :-

Chromite, or Chrome ironstone, FeCr₂O₄. Crocoisite, PbCrO₄.

GROUP III.B.—ZINC GROUP.

Zinc :--

Zincite, or Red zinc ore, ZnO.

Zinc blende, ZnS.

Calamine, ZnCO₃.

Silicious (electric) calamine, or Hemimorphite, 2ZnSiO₄.H₂O.

• Franklinite, (Zn,Fe)O.Fe₂O₃.

Manganese:-

Pyrolusite, MnO₂.

Braunite, Mn₂O₃.

Hausmannite, Mn₃O₄.

Manganite, Mn₂H₂O₄.

Psilomelane, (Mn,Ba)O.MnO₂.

Wad, mainly hydrated MnO₂ and Mn₂O₃.

Dialogite, or Manganese spar, MnCO₃.

Nickel:-

Copper-nickel, NiAs.

Chloanthite, NiAs₂.

Garnierite, Hydrated Ni, Mg silicates.

Cobalt :--

Smaltite, CoAs₂.

Cobaltite, CoAsS.

GROUP IV.—BARIUM GROUP.

Barium :-

Barytes, or Heavy spar, BaSO₄.

Witherite, BaCO₃.

Strontium :--

Celestine, SrSO₄.

Strontianite, SrCO₃.

Calcium :--

Calcite or Calcspar, Arragonite, Marble, Chalk, Limestone, CaCO₃.

Dolomite, or Magnesian limestone, (Ca,Mg)CO₃.

Gypsum, Selenite, or Alabaster, CaSO₄.2H₂O.

Anhydrite, CaSO₄.

Fluorspar, CaF2,

Fluor-apatite, 3Ca₃(PO₄)₂.CaF₂.

Chlor-apatite, 3Ca₃(PO₄)₂.CaCl₂.

GROUP V.—POTASSIUM AND MAGNESIUM GROUP.

Potassium :--

Nitre, or Saltpetre, KNO₃.

Sylvine, KCl.

Kainite (see Magnesium).

Carnallite, KCl.MgCl₂.6H₂O.

Alumstone, K₂O.3Al₂O₃.4SO₃.6H₂O.

Orthoclase (Potash) Felspar, K2O.Al2O3.6SiO2.

Sodium :--

Rocksalt, NaCl.

Natron, or Native soda, Na₂CO₃.10H₂O.

Trona, Na₂CO₃.NaHCO₃.3H₂O.

Soda nitre, or Chili saltpetre, NaNO₃.

Glauber salt, Na₂SO₄.10H₂O.

Glauberite, Na 2SO4. CaSO4.

Cryolite, AlF₃.3NaF.

Albite (Soda) Felspar, Na₂O.Al₂O₃.6SiO₂.

Borax, or Tincal, Na₂B₄O₇.10H₂O.

Magnesium :-

Periclase, MgO.

Brucite, MgO.H₂O.

Magnesite, MgCO₃.

Dolomite, or Magnesian limestone, (Mg,Ca)CO₃.

Spinel, MgO.Al₂O₃.

Kainite, MgSO₄.KCl.3H₂O.

Kieserite, MgSO₄.H₂O.

Epsomite, MgSO₄.7H₂O.

Boracite, 6MgO.MgCl₂.8B₂O₃.

Meerschaum, Mg₂H₂3SiO₂.H₂O.

Silica :--

Quartz, Rock crystal, Flint, SiO₂,

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